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(4) New amines, their use and preparation.

(57) Novel 3,3-diphenylpropylamines of formula I

anticholinergic drug, pharmaceutical compositions containing the novel amines, and methods for preparing the same.

wherein R1 signifies hydrogen or methyl, R2, R3 and R4 independently signify hydrogen, methyl, methoxy, hydroxy, carbamoyl, sulphanoyl or halogen, and X represents a tertiary amino group -NR5,R6, wherein R5 and R8 signify non-aromatic hydrocarbyl groups, which may be the same or different and which together contain at least three carbon atoms, and which may form a ring together with the amine nitrogen, their salts with physiologically acceptable acids and, when the compounds can be in the form of optical isomers, the racemic mixture and the individual enantiomers, their use as drugs, especially as anticholinergic agents, their use for preparing an

#### Description

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The present invention relates to novel 3,3-diphenylpropylamino derivatives, to pharmaceutical compositions containing the same, and to the use of said derivatives for preparing drugs.

Swedish patent No. 215 499 discloses certain 3,3-diphenylpropylyamines having an advantageous effect on the heart and circulation. These pharmacologically active 3,3-diphenylpropylamines are secondary amines. Said Swedish patent also discloses certain chemical intermediates which are tertiary amines carrying aromatic substituents on the amine nitrogen. Neither the end products (secondary amines) nor the intermediates (tertiary amines) have any hydroxy or methoxy groups as substituents in the ortho positions of the phenyl rings, but only meta and para substituents are specifically disclosed.

It is known that terodiline, a commercially available drug having the chemical formula

has anti-cholinergic properties, and is well resorbed in the body. However, this drug has a very long biological half-life and it is a multi-effect drug also having other pharmacological properties such as Ca-antagonist, noradrenaline antagonist and anti-histamine properties as well as a pronounced effect on the heart.

US-A-3.446.901, GB-A-1.169.944 and GB-A-1.169.945 disclose certain 3,3-diphenylpropylamine derivatives and pharmaceutical compositions having antidepressant activity, i.a. N,N-dimethyl-3-(2-methoxyphenyl)-3-phenylpropylamine, which is considered to be the closest prior art as regards chemical structure (see also the comparative tests reported at the end of this specification). DK-A-111.894 discloses a special process for preparing certain diphenylalkylamines having an effect on the heart and circulation. The specifically described compounds are primary or secondary amines, and none of them has any hydroxy or alkoxy substituent in ortho position of the phenyl rings. C.A. Vol. 97 (1982) 120105n discloses certain N-arylalkylisoquinolines which may have a hydroxy substituent in the ortho position of a phenyl ring. These compounds have sympatholytic activity and carry aromatic substituents on the nitrogen atom.

It is an object of the present invention to provide a novel class of 3,3-diphenylpropylamines having improved anti-cholinergic properties, especially in relation to the effects on these other systems and acute toxicity. In a first aspect the invention provides novel 3,3-diphenylpropylamines of formula I

$$R^2$$

$$O-OR^1$$

$$CH-CH_2-CH_2-X$$

$$R^3$$

wherein R¹ signifies hydrogen or methyl, R², R³ and R⁴ independently signify hydrogen, methyl, methoxy, hydroxy, carbamoyl, sulphanoyl or halogen, and X represents a tertiary amino group of formula II

wherein R<sup>5</sup> and R<sup>6</sup> signify non-aromatic hydrocarbol groups, which may be the same or different and which together contain at least three carbon atoms, preferably at least four carbon atoms, especially at least five carbon atoms, and wherein R<sup>5</sup> and R<sup>6</sup> may form a ring together with the amine nitrogen, said ring preferably having no other hetero atom than the amine nitrogen.

The compounds of formula I can form salts with physiologically acceptable acids, organic and inorganic, and the invention comprises the free bases as well as the salts thereof. Examples of such acid addition salts

include the hydrochloride, hydrobromide, hydrogen fumarate, and the like.

When the novel compounds can be in the form of optical isomers, the invention comprises the racemic mixture as well as the individual enantiomers as such.

A preferred sub-class of compounds according to the invention comprises tertiary amines of formula I, wherein each of R<sup>5</sup> and R<sup>6</sup> independently signifies C<sub>1-8</sub>-alkyl, especially C<sub>1-6</sub>-alkyl, or adamantyl, R<sup>5</sup> and R<sup>6</sup> together comprising at least three, preferably at least four carbon atoms. R5 and R6 may carry one or more hydroxy groups, and they may be joined to form a ring together with the amine nitrogen atom.

Presently preferred tertiary amino-groups X in formula I include the following groups a) - f), each of which may carry one or more hydroxy groups.

a) 
$$-N = \frac{CH(CH_3)_2}{CH(CH_3)_2}$$
, b)  $-N = \frac{CH_3}{C(CH_3)_3}$ , c)  $-N = \frac{CH_3}{C(CH_3)_2CH_2CH_3}$ ,

The following are examples of presently preferred specific compounds of formula 1:

N,N-diisopropyl-3-(2-hydroxy-5-methylphenyl)-3-phenylpropylamine and its (+)-isomer,

N-methyl-N-tert.butyl-3-(2-hydroxyphenyl)-3-phenylpropylamine,

N-methyl-N-tert.butyl-3-(2,4-dihydroxyphenyl)-3-phenylpropylamine,

N-methyl-N-tert.butyl-3,3-bis-(2-hydroxyphenyl)propylamine,

N,N-diisopropyl-3,3-bis-(2-hydroxyphenyl)propylamine,

N,N-diisopropyl-3-(2,5-dihydroxyphenyl)-3-phenylpropylamine,

N-methyl-N-tert.butyl-3-(2,5-dihydroxyphenyl)-3-phenylpropylamine,

N,N-diisopropyl-3-(2-methoxyphenyl)-3-phenylpropylamine,

N-[3-(2-methoxyphenyl)-3-phenylpropyl]-2,2,6,6-tetramethylpiperidine

In a second aspect of the invention provides methods for preparing the compounds of formula I, especially the following methods:

a) reacting a reactively esterified 3,3-diphenylpropanol of formula III

wherein R1-R4 are as defined above, and any hydroxy groups may be protected such as by methylation or benzylation, and wherein Y is a leaving group, preferably halogen or an alkyl or arylsulphonyloxy group, with an amine of formula IV

H-X IV

wherein X is as defined above, or

b) reducing a 3,3-diphenylpropionamide of formula V

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$$R^2$$

$$O-OR^1$$

$$CH-CH_2-CO-X$$

$$R^3$$

$$O-R^4$$

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wherein R1-R4 and X are as defined above and any hydroxy groups may be protected, preferably using a complex metal hydride,

c) N-methylating a secondary 3,3-diphenylpropylamine VI

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$$R^{2}$$

$$CH-CH_{2}-CH_{2}-NH-Z$$

$$VI$$

$$R^{3}$$

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wherein R1-R4 are as defined above and any hydroxy groups may be protected, and wherein Z has the same meaning as R5 and R6 with the exception of methyl, Z preferably being a hydrocarbyl group comprising at least three carbon atoms, the N-methylation preferably being carried out using formaldehyde or formic acid, or

d) reducing a 3,3-diphenylpropylamine of formula VIIa or VIIb

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VIIa

VIIb

wherein R<sup>1</sup>-R<sup>4</sup> and X are as defined above and any hydroxy groups may be protected, and W signifies a hydroxy group or a halogen atom, preferably by means of catalytic hydrogenation, and

i) when necessary splitting off hydroxy protecting groups in the compounds obtained, if desired after mono or di-halogenation of one or both of the phenyl rings, and/or

ii) if desired converting obtained bases of formula I into salts thereof with physiologically acceptable acids, or vice versa, and/or

iii) if desired separating an obtained mixture of optical isomers into the individual enantiomers, and/or

iv) if desired methylating an ortho-hydroxy group in an obtained compound of formula I, wherein R<sup>1</sup> is hydrogen and/or R<sup>4</sup> is hydroxy.

The above general methods can be carried out in a manner known per se and/or in accordance with the working examples described below, with due consideration of the desired amino groups and the substituents on the benzene rings.

on the benzene rings.

The removal of hydroxy protecting groups according to i) above can e.g. be done by treatment with hydrobromic acid, borontribromide or by catalytic hydrogenation.

The separation of mixtures of optical isomers, according to ii) above, into the individual enantiomers can e.g. be achieved by fractional crystallization of salts with chiral acids or by chromatographic separation on chiral columns.

Novel compounds of formula VIII

wherein R1-R4 are as defined above, and the corresponding protected compounds (e.g. comprising protected hydroxy groups), are useful as chemical intermediates for the preparation of e.g. the compounds of formula I, and they can be prepared by means of several different methods which are known per se, such as by addition of ethylene oxide (X) to a correspondingly substituted diphenylmethane (IX) in the presence of a suitable base such as sodium amide:

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The compounds VIII can also be prepared by reduction of the corresponding 3,3-diphenylpropionic acids, preferably using complex metal hydrides.

The 3,3-diphenylpropanols VIII can conveniently be converted into the corresponding reactively esterified derivatives III in a manner known per se by displacing the hydroxy groups with e.g. a halogen atom or an alkyl or arylsulphonyloxy group.

The 3,3-diphenylamides of formula V used as starting materials in method b), can e.g. be prepared by reacting the above mentioned 3,3-diphenylpropionic acids with an appropriate amine.

The secondary amines used as starting materials in method c) can conveniently be prepared by reacting a primary amine H<sub>2</sub>N-Z (wherein Z is as defined above) with a corresponding reactively esterified 3,3-diphenylpropanol in analogy with method a) above, or by reduction of the corresponding secondary 3,3-diphenylpropionamides in analogy with method b) above. The secondary amines can also be prepared by reduction of unsaturated hydroxyamines XI

$$R^2$$
O-OR<sup>1</sup>
C-CH<sub>2</sub>-CH=N-Z
XI
 $R^3$ 
O-R<sup>4</sup>

wherein R1-R4 and Z are as defined above, either in one step by catalytic hydrogenation, or by reduction to the corresponding saturated hydroxyamine, preferably using a complex metal hydride such as lithium aluminium hydride, followed by removal of the hydroxy group by catalytic reduction. As an alternative, the hydroxy group may first be split off as water, followed by reduction of the formed unsaturated amine.

The unsaturated hydroxy amines XI can conveniently be prepared by the addition of a Schiff base of formula XII

CH<sub>3</sub>-CH = N-Z XII wherein Z is as defined above, to a benzophenone of formula XIII

$$\begin{array}{c|c}
 & R^2 \\
 & \bigcirc -OR^1 \\
 & \bigcirc -OR^4
\end{array}$$
XIII

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wherein R1-R4 are as defined above, in the presence of a base, preferably a lithium organic base such as lithium diisopropylamide.

Also the starting materials VIIa, VIIb for process d) can be prepared by methods known per se, such as by addition of an organometallic compound XIVa or XIVb

to a ketoamine XVa or XVb respectively to form a corresponding hydroxy amine XVI

and, if desired, splitting off water from compound XVI.

In formulae XIVa, XIVb, XVa, XVb, XVI, R<sup>1</sup>-R<sup>4</sup> are as defined above, and Me signifies a metal such as magnesium or lithium.

In accordance with the invention the compounds of formula I, in the form of free bases or salts with physiologically acceptable acids, can be brought into suitable galenic forms, such as compositions for oral use, for injection, or the like, in accordance with accepted pharmaceutical procedures. Such pharmaceuti cal compositions according to the invention comprise the compounds of formula I in association with compatible pharmaceutically acceptable carrier materials, or diluents, as is well known in the art. The carriers may be any inert material, organic or inorganic, suitable for enteral, percutaneous or parenteral administration such as: water, gelatin, gum arabicum, lactose, microcrystalline cellulose, starch, sodium starch glycolate, calcium hydrogen phosphate, magnesium stearate, talcum, colloidal silicon dioxide, and the like. Such compositions may also contain other pharmaceutically active agents, and conventional additives such as stabilizers, wetting agents, emulsifiers, flavouring agents, buffers, and the like.

The compositions according to the invention can e.g. be made up in solid or liquid form for oral administration, such as tablets, capsules, powders, syrups, elixirs and the like, in the form of sterile solutions, suspensions or emulsions for parenteral administration, and the like.

The compounds and compositions according to the invention can be used for treating cholin-mediated disorders such as urinary incontinence. As is well known, the dosage depends on several factors such as the potency of the selected specific compound, the mode of administration, the age and weight of the patient, the severity of the condition to be treated, and the like. The daily dosage may, for example, be from about 0.05 mg

to about 4 mg per kilo of body weight, administered in one or more doses, e.g. containing from about 0.05 to about 200 mg each.

The invention will be further illustrated by the following non-limiting examples.

5 General 1H-NMR spectra were run in CDCl3 using a JEOL PMX60 spectrometer. In some cases, only a limited number of spectral peaks, useful for characterisation purposes, are reported. Reported yields mostly refer to crude material of sufficient purity to be taken to the next stage. 10 Solvents are abbreviated as follows: IPE = diisopropyl ether PET = petroleum ether Ether = diethyl ether 15 Amines are abbreviated as follows: IPA = diisopropyl amine TBA = tert.butyl amine Melting points were taken on a Koefler bench. Temperatures are in °C. 20 Water is used for the washing steps, unless otherwise stated. Example 1 25 Preparation of 4-phenyl-3,4-dihydrocoumarins a) 4-(2-Methoxy-5-methylphenyl)-6-methyl-3,4-dihydrocoumarin (I) 30 A mixture consisting of 2-methoxy-5-methylcinnamic acid (96.0 g, 0.5 mol), p-cresol (108 g, 1.0 mol), tetraline (200 ml), and conc. sulphuric acid (20 g) was heated slowly to refluxing temperature (145-150°). After 1 1/2 - 2 h, the mixture was cooled, taken up in ether, washed with water and sodium carbonate, dried and evaporated, giving 138 g (97%) crude oil. Two recrystallisations from acetone gave white crystals of the 35 desired lactone, m.p. 126-127°. 17.00 6.43 O C 76.57 C<sub>18</sub>H<sub>18</sub>O<sub>3</sub>(282.3)requires: 6.44 17.0 76.9 Found 40 b) 6-Hydroxy-4-phenyl-3,4-dihydrocoumarin (II) was prepared in a similar way in 97% yield from cinnamic acid and hydroquinone. M.p. 138° (IPE-Ether). 45 19.98 74.99 5.04 0 C C<sub>15</sub>H<sub>12</sub>O<sub>3</sub>(240.3)requires: 5.00 19.6 75.0 Found c) 4-(2-methoxy-4-methylphenyl)-7-methyl-3,4-dihydrocoumarin 50 was obtained in a similar way from 2-methoxy-4-methylcinnamic acid and m-cresol in 58% yield. M.p. 147-148° (IPE-acetone). 76.57 Н 6.43 О 17.00 С C<sub>18</sub>H<sub>18</sub>O<sub>3</sub>(282.3)requires: 55 6.31 17.2 76.4 Found The above lactone (90.g, 0.32 mol) in methylene chloride (500 ml) was refluxed with BBr<sub>3</sub> (115 g, 0.46 mol) for 24 h, the solution was concentrated, the residue was taken up in ether, the solution was washed with sodium carbonate and water, dried and evaporated, giving 80 g (93%) of a syrup which crystallized on 60 standing. Crystallization from IPE-PET gave white crystals of d) 4-(2-hydroxy-4-methylphenyl)-7-methyl-3,4-dihydrocoumarin (III), m.p. 137°.

	C <sub>17</sub> H <sub>16</sub> O <sub>3</sub> (268.3)requires: Found	С	76.10 76.2	н	6.01 6.30	0	17.89 17.0
5	e) 8-Hydroxy-4-phenyl-3,4-dihydro was obtained in a similar way from	coumarin (IV) m cinnamic a	cid and cat	echol in 18% y	yield. M.p	o. 136° (IPE).	
10	C <sub>15</sub> H <sub>12</sub> O <sub>3</sub> (240.2)requires: Found	С	74.99 75.0	<b>H</b>	5.04 5.01	0	19.98 19.9
15	f) 4-(2-Methoxyphenyl)-3,4-dihydrowas obtained in a similar way in 45 mixture was contaminated with methis by-product with ice-cold NaOH the next step.	% yield from thyl 3-(4-hydr	methyl 2-m oxyphenyl)-	3-(2-methoxyph	nenyl)-pro	pionate. After r	emoval of
20			Example 2				
	14						
25	Prepa	aration of 3,3	-diphenylpro	opionic acid es	ters		
30	a) Methyl 3-(2-methoxy-4-methylphoto-7-Methyl-4-phenyl-3,4-dihydrocomethyl iodide (100 g, 0.7 mol) and evaporated. The residue was dissogiving 86 g (92%) of a viscous of NMR: $\delta$ 6.6-7.2 (m 8H), 4.9 (t 1H)	oumarin (78 g, K₂CO₃ (55 g, olved in ether il.	0.327 mol) i , 0.4 mol) wa , the solution	n 150 ml metha as refluxed for t on was washed	24 h, filte with wate	red, and the so	lvent was
35	b) Methyl 3,3-bis-(2-methoxyphen was obtained in the same way in	yl)-propionate 96% yield fr	e (VII) om the lact	one (V) of Exa	mple 1f),	m.p. 84-87° (i	PE).
40	C <sub>18</sub> H <sub>20</sub> O <sub>4</sub> (300.4)requires: Found	С	71.98 71.4	Н	6.71 6.67	0	21.3 21.6
45	c) Methyl 3-(2,3-dibenzyloxypheny was obtained in a similar way in que methanol. In addition to K <sub>2</sub> CO <sub>3</sub> the	antitative yiel	d from the l	actone (IV) of E	Example 1 ne Nal. M	e) and benzyl o .p. 72° (IPE).	chloride in
	C <sub>20</sub> H <sub>28</sub> O <sub>4</sub> (452.5)requires: Found	С	79.63 79.9	Н .	6.24 6.15	0	14.14 14.1
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55	d) Methyl 3-(2-benzyloxyphenyl)-3 was obtained in a similar way as a chloride. NMR: $\delta$ 7.2 (m 14H), 4.9 (s 2H, t	a viscous oil	in 81% yiel		ri-3,4-dihy	drocoumarin a	nd benzyl
	e) Methyl 3-(2-methoxy-5-methylp was obtained in a similar way fron NMR: $\delta$ 7.4 (m 8H), 5.0 (t 1H), 3.	m 6-methyl-4	-phenyl-3,4-	dihydrocoumar	in in 96% 3H).	o yield.	
60	f) Methyl 3,3-bis-(2-methoxy-5-methoxy obtained in a similar way in NMR: $\delta$ 6.6-7.1 (m 6H), 5.1 (t 1H	quantitative y	ield from th	ne lactone (I) o		e 1a) and meth	ıyl iodide.
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g) Methyl 3-(2,5-dibenzyloxyphenyl)-3-phenylpropionate (XII) was obtained in a similar way in 90% yield from the lactone (II) of Example 1b) and benzyl chloride. NMR: $\delta$ 6.8-7.4 (m 18H), 5.0 (s 4H, t 1H), 3.7 (s 3H), 3.1 (d 2H).													
h) Methyl 3,3-bis-(2-benzyloxy-4-methylphenyl)propionate (XIII) was obtained in a similar way in 95% yield from the lactone (III) of Example 1d) and benzyl chloride. By GLC the product is homogenous, and by MS it has the correct M.W.	5												
<ul> <li>i) Ethyl 3-(2,4-dimethoxyphenyl)-3-phenylpropionate (XIV)         A mixture of ethyl cinnamate (88 g, 0.5 mol), dimethyl resorcinol (276 g, 2.0 mol) and conc. sulphuric acid (50 g) was stirred on a boiling water-bath for 2 h, whereafter all the volatile material was distilled off in vacuum. The residual oil was dissolved in ether, the solution was washed with sodium carbonate, dried, and evaporated giving 101 g (64%) of the title ester in the form of a viscous oil.         NMR: δ 6.4-7.2 (m 8H), 4.9 (t 1H), 4.0 (q 2H), 3.7 (s 6H), 3.0 (d 2H), 1.1 (t 3H).     </li> <li>j) Methyl 3,3-bis-(2,4-dimethoxyphenyl)propionate (XV)</li> </ul>													
j) Methyl 3,3-bis-(2,4-dimethoxyphenyl)propionate (XV) was obtained in a similar way from methyl 2,4-dimethoxycinnamate and dimethyl resorcinol. The product thus obtained contained about 23% of dimethyl resorcinol. It was taken to the next step without further purification.													
k) Methyl-3-(5-chloro-2-methoxyphenyl)-3-phenylpropionate 6-Chloro-4-phenyl-3,4-dihydrocoumarin (435 g, 1.68 mol. Preparation: T. Manimaran & V.T. Ramakrishnan, Ind. J. Chem. B 18 (1979) 328) is added to a hot solution of sodium hydroxide (140 g, 3.5 mol) in water (500 ml). The solution is chilled to 25°C and dimethyl sulphate (442 g, 3.5 mol) is added dropwise during 1 h with stirring and cooling at 25-35°C. The mixture is stirred for an additional 2 h whereupon a solution of 100 g of sodium													
The solution is chilled to 25°C and dimethyl sulphate (442 g, 3.5 mol) is added dropwise during 1 h with stirring and cooling at 25-35°C. The mixture is stirred for an additional 2 h whereupon a solution of 100 g of sodium hydroxide in 500 ml of water is added and the mixture is stirred until a clear solution is obtained. An excess of concentrated hydrochloric acid is added to precipitate the methoxy acid, which separates as an oil which slowly crystallizes. It is filtered off, washed with water and dried. Crystallization from 2-propanol gives colourless crystals of 3-(5-chloro-2-methoxyphenyl)-3-phenyl propionic acid, m.p. 144°C. Yield 455 g.													
The above acid (291 g, 1.0 mol) in 1 litre methanol containing 50 g concentrated sulphuric acid was refluxed for 8 h. The solvent was distilled off, the residue was taken up in ether, washed with water and sodium carbonat solution, dried and evaporated giving 300 g (100%) crude oil. Recrystallisation from IPE gave white crystals of the title compound, m.p. 65-66°.	30												
C <sub>17</sub> H <sub>17</sub> ClO <sub>3</sub> (304,8)requires: C 67.0 H 5.62 Cl 11.63 Found 68.1 5.82 11.7	35												
Example 3	40												
Preparation of 3,3-diphenylpropanols	45												
a) 3-(2-Methoxy-4-methylphenyl)-3-phenylpropanol (XVI)  The ester (VI) of Example 2a) (84 g, 0.295 mol) in 150 ml dry ether was added dropwise to a suspension of LiAlH4 (11.3 g, 0.295 mol) in 300 ml dry ether. The mixture was stirred overnight, then decomposed by the careful addition first of 11 g of water, then of 15% NaOH until a white granular precipitate was formed. The mixture was filtered, the filtrate was washed with water, dried, and evaporated giving 71 g (91%) of an oil which crystallized on standing. Recrystallization from IPE-PET gave white crystals, m.p. 83°.	50												
C <sub>17</sub> H <sub>20</sub> O <sub>2</sub> (256.4)requires: C 79.65 H 7.88 O 12.48 Found 79.4 7.89 12.7	55												
b) 3,3-Bis-(2-methoxyphenyl)propanol (XVII) was obtained in a similar manner in quantitative yield as a viscous oil from the ester (VII) of Example 2b).	60												
c) 3-(2,3-Dibenzyloxyphenyl)-3-phenylpropanol (XVIII) was obtained in a similar way as a viscous oil in 96% yield from the ester (VIII) of Example 2c).													
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d) 3-2(Benzyloxyphenyl)-3-phenylpropanol (XIX) was obtained in a similar way as an oil in 78% yield from the ester (IX) of Example 2d). e) 3-(2-Methoxy-5-methylphenyl)-3-phenylpropanol (XX) was obtained in a similar way as an oil in quantitative yield from the ester (X) of Example 2e). 5 NMR:  $\delta$  6.8-7.4 (m 7H), 4.7 (t 1H), 3.8 (s 3H), 3.7 (m 2H), 2.3 (s 3H), 2.0-2.3 (m 2H). 1) 3,3-Bis-(2-methoxy-5-methylphenyl)propanol (XXI) was obtained in a similar way in 98% yield from the ester (XI) of Example 2f). M.p. 89° (IPE). 10 8.05 15.98 75.97 Н 0 C<sub>19</sub>H<sub>24</sub>O<sub>3</sub>(300.4)requires: 75.9 8.02 16.1 Found 15 g) 3-(2,5-Dibenzyloxyphenyl)-3-phenylpropanol (XXII) was obtained in a similar way in 88% yield from the ester (XII) of Example 2g). M.p. 78° (IPE). 82.05 6.65 11.31 C<sub>29</sub>H<sub>28</sub>O<sub>3</sub>(424.5)requires: 20 6.62 Found 82.0 11.2 h) 3,3-Bis-(2-benzyloxy-4-methylphenyl)propanol (XXIII) was obtained in a similar way as an oil in 93% yield from the ester (XIII) of Example 2h). 25 i) 3-(2,4-Dimethoxyphenyl)-3-phenylpropanol (XXIV) was obtained as a golden oil in 92% yield from the ester (XIV) of Example 2i). NMR:  $\delta$  6.5-7.2 (m 8H), 4.5 (t 1H), 3.8 (s 6H), 3.6 (m 2H), 2.0-2.6 (m 3H). 30 j) 3,3-Bis-(2,4-dimethoxyphenyl)propanol (XXV) was obtained in a similar way from the impure ester (XV) of Example 2j). By NMR, the product contains about 20% of dimethyl resorcinol. k) 3-(4-Fluorphenyl)-3-(2-methoxyphenyl)propanol (XXVI) A Grignard reagent was prepared in the usual manner from o-bromoanisole (93.5 g, 0.5 mol) and magnesium (12 g, 0.5 mol) in 100 ml dry ether. A solution of p-fluorobenzaldehyde (62 g, 0.5 mol) in 100 ml ether was added dropwise to this solution. After about 1 h, the mixture was decomposed with NH<sub>4</sub>Cl and worked up, giving 100.6 g (87%) of 4-fluoro-2'-methoxy-diphenylmethanol. Recrystallization from IPE-PET gave white crystals, 40 m.p. 88°. 5.64 C<sub>14</sub>H<sub>13</sub>FO<sub>2</sub>(232.3)requires: C 72 40 72.9 5.75 Found 45 The obtained carbinol (46.2 g, 0.2 mol) in 600 ml ethanol was hydrogenated in the presence of 4 g of 5% Pd/C catalyst. After about 5-6 h, the reaction was complete and the mixture was worked up giving 40 g (93%) of 4-fluoro-2'-methoxy-diphenylmethane as a clear oil. NMR: 6.8-7.2 (m 8H), 4.0 (s 2H), 3.8 (s 3H). The obtained methane derivative (71 g, 0.33 mol) in 100 ml ether was added to a solution of NaNH2 prepared in situ from sodium (8.5 g, 0.37 mol) in about 300 ml of NH<sub>3</sub>. After about 1 h, a solution of ethylene oxide (17.5 g, 0.395 mol) in 75 ml ether was added dropwise. The mixture was stirred for 2 h, and most of the ammonia was then removed with a stream of air. Solid NH<sub>4</sub>Cl was then added, followed by the addition of water. The organic phase was separated, washed with water and 2N HCI, dried and evaporated, giving 81.5 g (95%) of the title 55 compound. M.p. 61° (IPE-PET). 6.58 73.82 н C<sub>16</sub>H<sub>17</sub>FO<sub>2</sub>(260.3)requires:

!) 3-(5-Chloro-2-methoxyphenyl)-3-phenylpropanol

Found

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The ester from Example 2k) (91.5 g, 0.3 mol) in 500 ml dry ether was added dropwise under nitrogen to LiAlH<sub>4</sub> (11.4 g, 0.3 mol) in 200 ml dry ether. The mixture was stirred at room temperature overnight, then decomposed with 11 g water and 11 g 15% NaOH solution. Work up gave 72.5 g (87.5%) colourless oil.

74.1

6.77

Recrystallization from IPE gave white crystals of the title compound, m.p. 80°. 12.81 6.19 CI 69.43 н C<sub>16</sub>H<sub>17</sub>ClO<sub>2</sub>(276.8)requires: 6.44 12.9 70.1 Found 5 Example 4 10 Preparation of 3,3-diphenylpropyl-p-toluene sulphonates 15 a) 3,3-Bis-(2-methoxyphenyl)propyl-p-toluene sulphonate (XXVII) The propanol (XVII) of Example 3b) (35 g, 0.128 mol) in 100 ml chloroform containing 30 ml pyridine was cooled to about -10° and then treated with p-toluene sulphonyl chloride (29 g, 0.15 mol). After standing in the cooler (about +5°C) overnight, the mixture was poured into ice-water, the organic phase was washed with water and cold 2N HCl, dried, and the solvent was distilled off at < 50°C, giving a crude oil in quantitative yield. 20 Recrystallization from IPE gave white crystals of low and indefinite m.p. 7.52 6.14 S C<sub>24</sub>H<sub>26</sub>O<sub>5</sub>S(426.5)requires: 67.58 6.22 7.76 66.8 Found 25 b) 3-(2-Methoxy-4-methylphenyl)-3-phenylpropyl-p-toluene sulphonate (XXXI) was obtained in quantitative yield from the propanol (XVI) of Example 3a). 30 c) 3-(2,3-Dibenzyloxyphenyl)-3-phenylpropyl-p-toluene sulphonate (XXVIII) was obtained in a similar way as a thick oil in 88% yield from the propanol (XVIII) of Example 3c). d) 3-(2-Benzyloxyphenyl)-3-phenylpropyl-p-toluene sulphonate (XXIX) was obtained in i similar way in 98% yield from the propanol (XIX) of Example 3d). 35 e) 3-(2-Methoxy-5-methylphenyl)-3-phenylpropyl-p-toluene sulphonate (XXX) was obtained in quantitative yield from the propanol (XX) of Example 3e). M.p. 64° (IPE-PET). , 8.09 40 S 69.67 6.10 C<sub>23</sub>H<sub>24</sub>O<sub>4</sub>S(396.5)requires: 7.85 6.20 69.8 Found 45 f) 3,3-Bis-(2-methoxy-5-methylphenyl)-propyl-p-toluene sulphonate (XXXII) was obtained in quantitative yield from the propanol (XXI) of Example 3f). M.p. 117° (acetone-PET). 68.7 6.65 S 7.05 C<sub>26</sub>H<sub>30</sub>O<sub>5</sub>S(454.5)requires: 68.8 6.66 7.11 Found 50 g) 3-(2,5-Dibenzyloxyphenyl)-3-phenylpropyl-p-toluene sulphonate (XXXIII) was obtained in a similar manner in quantitative yield from the propanol (XXII) of Example 3g). 55 h) 3,3-Bis-(2-benzyloxy-4-methylphenyl)-propyl-p-toluene sulphonate (XXXIV) was obtained in a similar way in 86% yield from the propanol (XXIII) of Example 3h). i) 3-(2,4-Dimethoxyphenyl)-3-phenylpropyl-p-toluene sulphonate (XXXV) was in the same way obtained in 96% yield from the propanol (XXIV) of Example 3i). 60 j) 3,3-Bis-(2,4-dimethoxyphenyl)-propyl-p-toluene sulphonate (XXXVI) was obtained in the same manner from the propanol (XXV) of Example 3j). The product was contaminated with dimethyl resorcinol. 65

	k) 3-(4-Fluorphenyl)-3-(2-methowas obtained in a similar way in	xyphenyl)- n 88% yie	propyl-p-tolue	ne sulphoni ropanol (XX	ate (XXXVII) VI) of Examp	ple 3k). N	М.р. 67°	(IPE).
5	$C_{23}H_{23}FO_4S(414.5)$ requires: Found	С	66.65 67.1	н	5.59 5.69	S		7.74 7.78
10	l) 3-(2-Methoxyphenyl)-3-phenyl A mixture of anisole (1080 g, 1 refluxed for 2 h in an apparatus oily residue was dissolved in eth 304 g (77%) of a pale yellow oil, b	0 mol), be equipped v ner, washe o.p. 115-11	enzyl alcohol (2 with a water se d with water a 8°/0.4 Torr. By	16 g, 2 mol) parator. Exc and sodium of NMR, it is a	and p-toluer cess of aniso carbonate, dr 1:1 mixture of	ile was the ried and f of o-meth	ien distil fractiona oxy and	led off, the ted, giving p-methoxy
15	diphenyl methane. This material ethylene oxide, as in the prepara converted as described above to isolated in 35% yield after two	was conve tion of the o a mixture	erted to a mixt propanol (XXV e of p-toluene	ure of the c i) of Exampl sulphonates	orresponding e 3k). This mi from which t	propand exture of p	ols by rea propanol	action with s was then
20	C <sub>23</sub> H <sub>24</sub> O <sub>4</sub> S(396.5)requires: Found	С	69.67 69.3	н	6.10 6.00	S		8.09 8.17
25	m) 3-(5-Chloro-2-methoxypheny The alcohol from Example 3l) portionswise in the cold with p-to h, solvent was evaporated under N HCl, dried and evaporated giv title compound, m.p. 89-90°.	(66 g, 0.24 oluene-sul vacuum at	4 mol) in 300 $\mu$ phonyl chlorid $t < 50^{\circ}$ , the re	ml chlorofor e (55 g, 0.29 sidue was ta	m containing mol). The mi ken up in eth	xture was ier, washi	s kept at ed with v	5°C for 18 vater and 2
30	C <sub>23</sub> H <sub>23</sub> ClO <sub>4</sub> S(430.96)requires: Found	С	64.10 H 64.4	5.38 5.45	0	7.44 7.04	CI	8.23 8.17
35		٠	Exampl	<u>e 5</u>				
40	<u>Pr</u>	eparation	of tertiary 3,3	-diphenylpro	ppylamines			
45	a) N,N-Diisopropyl-3,3-bis-(2-me) The tosylate (XXVII) of Exa diisopropylamine was heated in a residue was treated with excess extracted with 2N HCI. This extra dried, decoloured, filtered and e	mple 4a) a pressure of 2N NaC act was wa	(42.6 g, 0.1 bottle at 80° for the and extracted shed with eth	mol) in 100 or 4-6 days. Ved with ethe er, basified,	oml aceton Volatile mater r. The extract extracted with	itrile and ial was th was was th ether, v	en evap hed with washed	orated, the water and with water,
50	oxalic acid salt by treating an a M.p. 160-161° (acetone).	acetone so	olution of the	base with o	ne equivalen	t of oxal	ic acid i	n acetone.
55	$C_{25}H_{35}NO_6(445.6)$ requires: Found	С	67.39 H 67.2	7.92 8.22	N .	3.14 2.94	0	21.55 21.9

b) N,N-Diisopropyl-3-(2,3-dibenzyloxyphenyl)-3-phenylpropylamine (XXXIX)

The free base was obtained in the same way in 75% yield from the tosylate (XXVIII) of Example 4c). NMR: 6.9-7.2 (m 18H), 5.0 (s 4H), 0.9 (d 12H).

c) N,N-Diisopropyl-3-(2-methoxy-5-methylphenyl)-3-phenylpropylamine (XL), hydrogenfumarate The free base was obtained in 69% yield from the tosylate (XXX) of Example 4e). It was converted to the fumaric acid salt in the usual manner. M.p. 176° (acetone).

60

C <sub>27</sub> H <sub>37</sub> NO <sub>5</sub> (455.7)requires: Found	С	71.17 71.3	н	8.20 8.27	N	3.07 3.04	0	17.6 17.9	
d) N-N-Diisopropyl-3-(2-methox The free base was obtained in m.p. 147-148° (acetone).	y-4-methy 25% yield	phenyl)-	3-phenyl e tosylate	propylami e (XXXI) o	ne (XLI) f Exampl	, hydrogenfi e 4b). The fi	ımarate ımaric a	cid salt had	5
C <sub>27</sub> H <sub>37</sub> NO <sub>5</sub> (455.7)requires: Found	С	71.17 71.3	н	8.20 8.14	N	3.07 3.00	0	17.6 17.6	10
e) N,N-Diisopropyl-3,3-bis-(2-m The free base was obtained i hydrochloride with ethereal HC	n 78% viel	d from th	ne tosvla	ite (XXXII)	of Exam	iple 4f). It wa	as conve	erted to the	15
C <sub>25</sub> H <sub>38</sub> NO <sub>2</sub> Cl(420.1)requires: Found		1.49 H 1.6	9.1 9.0		3.33 3.27	O 7.61 7.93	CI	8.44 8.36	20
f) N,N-Diisopropyl-3-(2,5-dibenz The free base was obtained NMR: 8 6.6-7.2 (m 18H), 5.0 (s	in 70% yi	eld from	the tos	ylate (XX)	(LIII) (III) of E	xample 4g).			25
g) N,N-Diisopropyl-3,3-bis-(2-bit) The free base was obtained NMR: δ 6.8-7.2 (m 16H), 4.8 (s	in 62% yi s 4H, t 1H	eld from ), 0.9 (d	the tos 12H).	ylate (XX)	KIV) of E	xample 4h)			30
h) N,N-Diisopropyl-3-(2,4-dimed The free base was obtained NMR: 6.5-7.3 (m 8H), 4.4 (t 1h i) N,N-Diisopropyl-3,3-bis-(2,4-dimed The free base was obtained NMR: δ 6.5-7.3 (m 6H), 4.6 (t	in 56% yi 1), 3.8 (s 6 dimethoxyr in 34% y	eld from SH), 1.0 ( chenyl)pr eld from	the tos (d 12H). opylaming the tos	ylate (XXX ne (XLVI) ylate (XXX	XV) of E				35
j) N,N-Diisopropyl-3-(4-fluoropl The free base was obtained	nenvi)-3-(2	-methox\	phenvl)	propylami	ne XLVII XVII) of	) Example 4k	).		40
k) N,N-Diisopropyl-3-(2-metho: The free base was obtained i fumaric acid salt in the usual	n 86% viel	d from th	e tosvlat	te (XLVIII)	of Exam	ple 4I) and w	as conv	erted to the	45
C <sub>26</sub> H <sub>36</sub> NO <sub>5</sub> (441.6)requires: Found	С	70.72 70.8	н	7.99 7.93	N	3.28 3.28	O.	18.12 18.1	50
i) N-[3-(2-Methoxyphenyl)-3-ph This compound was obtaine	d in the sa	me way	tetrame in 54%	thylpiperio	the (LXI	<u>V)</u> rlate (XLVIII)	of Exar	mple 4I) and	30
2,2,6,6-tetramethylpiperidine. N C <sub>25</sub> H <sub>35</sub> NO(365.6)requires: Found	И.р. 100° (	8	2.14 2.0	Н		9.65 N 9.62		3.83 3.57	55
m) N,N-diisopropyl-3-(5-chloro The tosylate from Example 4	n) (43.1 a.	0.1 mol) v	was heat	ed for 4 da	ays at 80°	° with diisop	ropylami	ine (50 g, 0.5	60
mol) in 100 ml acetonitrile, givin	g 23 g (64º	/o) of crue	de title c	ompound.	.By GC, i	t is at least 9	3% pure	<b>.</b> .	65

	n) N-[3-(2-Benzyloxyphenyl)-3-ph This compound was similarly pre- lidine. It was obtained as a sticky of (Example 9ab)).	epared fro	m the	tosylate	$\overline{(XXI)}$	() of Exa	ample	4d) and logue	d 2,2,5 withou	,5-tetra it furthe	imeth er puri	ylpyrro- ification
5	o) N-[3-(2-Benzyloxyphenyl)-3-phenylpropyl]-4-hydroxy-2,2,6,6-tetramethylpiperidine This compound was similarly prepared from the tosylate (XXIX) of Example 4d) and 4-hydroxy-2,2,6,6-tetramethylpiperidine, and It was obtained as a sticky oil which was converted to the hydroxy compound without further purification (Example 9ac)).											
10	p) N-(2-Hydroxy-1,1-dimethylethy This compound was similarly p panol. The solid product was cry	l)-3-(2-be	om the	e tosvlat	e (X)	(IX) of E	xamp	le 4d)	and 2⋅ 03°C. ∣	-amino- It was i	·2-me used	thylpro- as start
15	material in Example 7p).											
	C <sub>26</sub> H <sub>31</sub> NO <sub>2</sub> (389.5)requires: Found	С	80.1 80.0			8.02 8.09	N		3.60 3.69	0		8.22 8.51
20	q) N-(1-Adamantyl)-3-(2-benzylox This compound was similarly pr was used as start material in Exa melted at 225°C.	epared fro	m the	tosylate	(XX	IX) of E	xampl drate	e 4d) a was pr	nd 1-a epare	minoad d in ac	damar etonit	ntane. It rile and
<i>25</i> .	C <sub>32</sub> H <sub>37</sub> NO.HCl.1/2H <sub>2</sub> O(497.1)red Found	quires:	С	77.31 77.3	Н	7.91 8.23	N	2.82 2.65	0	4.83 5.04	ĊI	7.13 7.14
30						_						
			<u> </u>	Example	<u>6</u>							
	•											
35	Prepa	aration of	secon	ndary 3,3	-dipl	nenylpro	pylan	nines				
35 40	a) N-tert.Butyl-3,3-bis-(2-methox) The tosylate (XXVII) of Examp Example 5, giving the free base in M.p. 135-136° (acetone-ether).	yphenyl)pi le 4a) wa	ropylar s heat	mine (L) ted with	, hyc	lrogen (	oxalat	e tert.b	utylam id salt	nine as in the u	desc Isual I	ribed in manner.
	a) N-tert.Butyl-3,3-bis-(2-methox The tosylate (XXVII) of Examp Example 5, giving the free base in	yphenyl)pi le 4a) wa	ropylar s heat	mine (L) ted with th was c	, hyc	lrogen (	oxalat	e tert.b	utylam id salt 3.36 3.36	nine as in the u	desc usua <u>l</u> i	ribed in manner. 22.99 23.4
40	a) N-tert.Butyl-3,3-bis-(2-methox) The tosylate (XXVII) of Examp Example 5, giving the free base in M.p. 135-136° (acetone-ether).  C23H31NO6(417.5)requires:	yphenyl)pi le 4a) wa 178% yield C C /phenyl)-3	ropylars heat d, which 66.1 65.6	mine (L) ted with th was c  7 H	, hyc a lar onve	rogen of ge excepted to 1 7.48 7.31	oxalatess of the ox	e tert.b alic ac	3.36 3.36	in the u	ısua <u>l</u> ı	22.99 23.4
40 45	a) N-tert.Butyl-3,3-bis-(2-methox: The tosylate (XXVII) of Examp Example 5, giving the free base in M.p. 135-136° (acetone-ether).  C <sub>23</sub> H <sub>31</sub> NO <sub>6</sub> (417.5)requires: Found  b) N-tert.Butyl-3-(2,3-dibenzylox) The free base was obtained as	yphenyl)pi le 4a) wa 178% yield C C /phenyl)-3	ropylars heat d, which 66.1 65.6	mine (L) ted with th was c  7 H	, hyc a lar onve	7.48 7.31  (Li), hosylate (	oxalatess of the ox	e tert.b alic ac	3.36 3.36	in the u	ısua <u>l</u> ı	22.99 23.4
40 45 50	a) N-tert.Butyl-3,3-bis-(2-methox) The tosylate (XXVII) of Example 5, giving the free base in M.p. 135-136° (acetone-ether).  C <sub>23</sub> H <sub>31</sub> NO <sub>6</sub> (417.5)requires: Found  b) N-tert.Butyl-3-(2,3-dibenzylox) The free base was obtained as a m.p. 184-185° (acetone-methanom).  C <sub>33</sub> H <sub>38</sub> NO <sub>2</sub> CI(516.1)requires:	yphenyl)pi le 4a) wa 178% yield C (phenyl)-3 above in 76 bl-IPE)	ropylars heat d, which 66.1 65.6 -pheny 8% yie	mine (L) ted with th was c  7 H ylpropyla eld from t  76.79 76.3	, hyce a lar onve	7.48 7.31  e (Li), hosylate ( 7.42 7.30	oxalatess of the ox	e tert.b alic aci	3.36 3.36 ample	O	e HCl	22.99 23.4 salt had 6.87 6.81
40 45 50	a) N-tert.Butyl-3,3-bis-(2-methox) The tosylate (XXVII) of Examp Example 5, giving the free base in M.p. 135-136° (acetone-ether).  C23H31NO6(417.5)requires: Found  b) N-tert.Butyl-3-(2,3-dibenzylox) The free base was obtained as a m.p. 184-185° (acetone-methano  C33H38NO2CI(516.1)requires: Found  c) N-tert.Butyl-3-(2-benzyloxyphe The free base was obtained in	yphenyl)pi le 4a) wa 178% yield C (phenyl)-3 above in 76 bl-IPE)	ropylars heat d, which 66.1 65.6 -pheny 8% yiel	mine (L) ted with th was c  7 H ylpropyla eld from t  76.79 76.3	, hyce a lar onve	7.48 7.31 e (Li), h osylate ( 7.42 7.30 ), hydro(XXIX) o	oxalatess of the oxide o	e tert.b alic aci	3.36 3.36 ample	O	e HCI CI acid	22.99 23.4 salt had 6.87 6.81

d) N-tert.Butyl-3-(2-methoxy-5-methylphen The free base was obtained in 90% yield f HCl, it gave a somewhat hygroscopic salt (ethanol-ether).	rom th	ne tosylat	e (XX	(X) of E	Examp	ole 4e). V	Vhen	treated of wat	l with er. M	ethereal i.p. 171°	5
$C_{21}H_{29}NO.HCl.1/4H_2O(352.5)$ (requires): Found	С	71.55 71.8	Н	8.74 8.72	. <b>N</b>	3.97 4.05	0	5.67 5.57	CI	10.06 10.1	>
e) N-tert.Butyl-3-(2-methoxy-4-methylphen The free base was obtained in quantitation.p. 138-149° (methanol-isopropanol). It v	e yiel	d from th	e tos	ylate (	XXXI)	of Exan	oride nple 4	b). The	e HCI-	-salt had	10
C <sub>21</sub> H <sub>30</sub> NOCl.3/4H <sub>2</sub> O(361.5)requires: Found	(			-	8.80 8.76	N	3.8 3.9:			9.81 9.75	15
f) N-tert.Butyl-3,3-bis-(2-methoxy-5-methylp The free base was obtained in quantitativ m.p. 242° (acetone).	oheny e yiel	l)-propyla d from th	ımine e tos	(LV), ylate (2	hydro XXXII)	ochloride of Exar	<u>e</u> nple 4	4f). The	HCI-	salt had	20
C <sub>23</sub> H <sub>34</sub> NOCl(392.0)requires: Found	С	70.47 70.2	' Н		8.74 B.81	N	3.5 3.4			9.05 8.99	25
g) N-tert.Butyl-3-(2,5-dibenzyloxyphenyl)-3- The free base was obtained in 85% yiem.p. 188° (ethanol-ether).	-phen eld fro	ylpropyla om the to	mine sylat	(LVI), e (XXX	hydro (III) o	ochloride f Examp	e ole 4g	). The	HCI	salt had	30
C <sub>33</sub> H <sub>38</sub> NO <sub>2</sub> Cl(516.1)requires: Found	С	76.79 77.2	Н	7.42 7.50	N	2.71 2.64	0	6.20 6.53	CI	6.87 6.85	35
h) N-tert.Butyl-3,3-bis-(2-benzyloxy-4-meth The free base was obtained in 94% yie m.p. 210° (acetone-ether).	ylphe ld fro	nyl)-prop m the to	ylami sylate	ne (LV e (XXX	II), hy	ydrochlo f Examp	<u>ride</u> ole 4h	). The	HCL-	salt had	40
C <sub>35</sub> H <sub>42</sub> NO <sub>2</sub> Cl(544.2)requires: Found	С	77.25 77.6	Н	7.78 7.82	<b>N</b>	2.57 2.35	0	5.89 6.08	CI	6.52 6.55	45
i) N-tert.Butyl-3-(2,4-dimethoxyphenyl)-3-pi The free base was obtained in 84% yield f (acetone-ethanol-ether).	henylr rom th	oropylami ne tosylat	ne (L e (XX	VIII), h	nydro Exam	chloride iple 4i). T	Γhe Η	CI-salt	had m	n.p. 196°	50
C <sub>21</sub> H <sub>30</sub> NO <sub>2</sub> CI(363.9)requires: Found	С	69.31 69.3	н	8.31 8.44	N	3.85 3.80	0	8.79 8.89	CI	9.74 9.81	. 66
j) N-tert.Butyl-3,3-bis-(2,4-dimethoxypheny The free base was obtained in 60% yie m.p. 251° (methanol-acetone).	l)-pro eld fro	pylamine om the to	(LIX) osylat	, hydro le (XX	ochlo XVI) (	ride of Exam	ple 4	j). The	HCI-	salt had	55
C <sub>23</sub> H <sub>34</sub> NO <sub>4</sub> Cl(424.0)requires: Found	С	65.15 · 64.5	Н	8.08 8.06	N	3.30 3.57	0	15.09 15.3	CI	8.36 8.67	60

	k) N-tert.Butyl-3-(4-fluorophenyl)-3-(2-metl The free base was obtained in 89% yiem.p. 194° (ethanol-acetone).	noxypl eld fro	nenyl)-prop m the tos	oylar ylate	nine (LX), (XXXVII)	hydroch of Exam	loride ple 4k	). The	HCI-	salt had
5	C <sub>20</sub> H <sub>27</sub> NOFCI(351.9)requires: Found	С	68.26 68.9	Н	7.73 7.97	N	3.98 4.0			10.08 9.69
10	l) N-tert.Butyl-3-(2-methoxyphenyl)-3-phen The free base was obtained in 88% y m.p. 205°.	ylprop ield fr	oylamine (loom the to	_XI), sylat	hydrochlo e (XLVIII)	oride of Exam	npie 4l	). The	HCI-	salt had
15	C <sub>20</sub> H <sub>28</sub> NOCl(333.9) requires: Found	С	71.94 71.9	Н	8.45 8.44		4.20 4.67			4.79 4.79
20	m) N-(1,1-Dimethylpropyl)-3-(2-methoxy-5- The free base was obtained in 95% yield HCl-salt had m.p. 188-189° (ethanol-aceto	from	iphenyi)-3 the tosyla	-phe te (X	nylpropyla (XX) of Exa	mine (L) ample 4e	(II), hy ) and t	drochl tert. an	oride nylam	ine. The
25	C <sub>22</sub> H <sub>32</sub> NOCl(362.0)requires: Found	C	73.00 73.4	Н	8.91 N 8.98	3.87 3.83	0	4.42 4.61	CI	9.80 9.51
30	n) N-(1,1-Dimethylpropyl)-3,3-bis-(2-methor). The free base was obtained in 94% yield HCI-salt had m.p. 210° (ethanol-acetone).	from	nethylpher the tosylat	nyl)p e (X	ropylamine XXII) of Ex	e (LXIII), cample 4	hydro f) and	chlorid tert. ar	l <u>e</u> nylan	nine. The
	C <sub>24</sub> H <sub>36</sub> NO <sub>2</sub> Cl(406.0)requires: Found	C	71.00 71.1	Н	8.94 N 9.01	3.45 3.60	0	7.88 7.92	CI	8.73 8.73
35										
40	o) N-tert.Butyl-3-(5-chloro-2-methoxypher The tosylate from Example 4m) (43.1 g, 0 0.5 mol) and the mixture was heated in a p (100%) crude title compound. The bas hydrochloride salt, m.p. 216-218°.	.1 mol	) in 100 ml re bottle a	acet t 80°	tonitrile wa of for 4 day	s. The us	sual we	ork-up	affor	ded 32 a
45	C <sub>20</sub> H <sub>26</sub> CINO.HCI(368.36)requires: Found	C	65.21 65.1	Н	7.39 7.39	N N	3.8 3.9			19.25 18.7
										•
<i>50</i>			Example	7_	•					
	Preparation of tertiary 3,	3-dipt	envloropy	lamir	nes from s	secondar	y amir	nes		
	r roparation of to that y									
55 60	a) N-Methyl-N-tert.butyl-3-(2-methoxypher A mixture of the secondary amine (LXI) 37% formaldehyde solution (12.5 g, 0.12 m with NaOH, and extracted with ether. The (94%) of a crude oil. The HCl-salt was p	of Exa nol) wa extract	ample 6I) ( s refluxed t was wash	29.7 for 1 red v	g, 0.1 mol 18-24 h. Th vith water,	), formic e mixture dried an	acid ( e was f d evap	13.8 g, then co orated	ooled I, givii	, basified
	C <sub>21</sub> H <sub>30</sub> NOCl(347.9)requires: Found	С	72.49 71.9		8.69 8.79	9 N	4.0 4.2	3 CI		10.19 10.1
65										

b) N-Methyl-N-tert.butyl-3-(2-methoxy-5-methylphenyl)-3-phenylpropylamine (LXVI), hydrochloride  The free base was obtained in the same way in 89% yield from the amine (LIII) of Example 6d). The HCl-salt had m.p. 161° (acetone).												
C <sub>22</sub> H <sub>32</sub> NOCl(362.0)requires: Found	С	73.00 73.0	Н	8.91 8.96	N	3.87 3.94	0	4.42 4.59	CI	9.08 9.77	J	
c) N-Methyl-N-tert.butyl-3,3-bis-(2-methox The free base was obtained in 96% yield (acetone-ether).	yphen from t	yl)propylar he amine (	nine L) of	(LXVII f Exam	), hyo ple 6a	drochlor a). The H	ride ICI-sa	ılt had ı	m.p. 1	87-190°	10	
C <sub>22</sub> H <sub>33</sub> NOCl(378.0)requires: Found	С	69.91 69.9	Н	8.54 8.56	N	3.71 3.53	0	8.47 8.93	CI	9.38 8.92	15	
d) N-Methyl-N-tert.butyl-3-(2-methoxy-4-m The free base was obtained in 96% yie	ethylp ld fro	henyi)-3-p m the ami	heny ne (	ripropyl LIV) of	lamin Exa	e (LXVI mple 6e	<u>ll)</u> ). M.į	o. 64°	(IPE)		20	
C <sub>22</sub> H <sub>31</sub> NO(325.5)requires: Found	С	81.17 81.0	Н		).60 ).83	N	4.30 4.19			4.92 5.03	25	
e) N-Methyl-N-tert.butyl-3,3-bis-(2-methox The free base was obtained in 97% yie	y-5-me eld fro	ethylpheny m the ami	n)pro	pylami LV) of	ne (L Exar	_XIX) nple 6f)	. М.р	. 95° (	IPE).		<i>30</i>	
$C_{24}H_{35}NO_2(370.0)$ requires: Found	С	78.00 78.1	Н		9.55 9.57	N	3.79 3.79			8.66 8.80		
f) N-Methyl-N-tert.butyl-3-(4-fluorophenyl)- he free base was obtained in 82% yield (ethanol-acetone).	-3-(2-n from	nethoxyph the amine	enyl) (LX)	propyl of Ex	amino	e (LXX) e 6k). Ti	, hydr ne HC	ochlor 3-salt i	ide nad m	ı.p. 218°	35	
C <sub>21</sub> H <sub>29</sub> NOCIF(365.9)requires: Found	С	68.93 69.0	н		7.99 7.97	N	3.8 3.9			9.69 9.60	40	
g) N-(1,1-Dimethylpropyl)-N-methyl-3-(2-mydrochloride  The free base was obtained in 98% m.p. 176-177° (acetone).										salt had	45	
C <sub>23</sub> H <sub>34</sub> NOCI(376.0)requires: Found	С	73.47 73.4	Н		9.11 9.15	N	3.7 3.7			9.43 9.41	50	
h) N-(1,1-Dimethylpropyl)-N-methyl-3,3-bis The free base was obtained in 89% yield (acetone-ether).	s-(2-m I from	ethoxy-5-r the amine	neth (LXI	ylphen II) of E	yi)pro xamp	opylamin ole 6n). 1	ie (D The H	(XII), h CI-salt	ydroc had n	hloride n.p. 147°	55	
C <sub>25</sub> H <sub>37</sub> NO <sub>2</sub> Cl(420.1)requires: Found	C	71.49 70.8	Н	9.12 9.20	N	3.34 3.63	0	7.62 7.74	CI	8.44 8.42	60	
i) N-Methyl-N-tert.butyl-3-{2,4-dimethoxyp This compound was obtained as an oi	henyl) I in qu	-3-phenylp jantitative	oropy yield	ylamine I from	(LX) the a	XIII) amine (L	-VIII) -	of Exa	mple	6i).	<i>65</i>	

NMR: 6.5-7.3 (m 8H), 4.3 (t 1H), 3.8 (s 6H), 2.3 (s 3H), 1.0 (s 9H).

j) N-Methyl-N-tert.butyl-3-(2,5-dibenzyloxyphenyl)-3-phenylpropylamine (LXXIV)
This was obtained as an oil in 95% yield from the amine (LVI) of Example 6g).

k) N-Methyl-N-tert.butyl-3,3-bis-(2-benzyloxy-4-methylphenyl)propylamine (LXXV), hydrochloride
The free base was obtained in 92% yield from the amine (LVII) of Example 6k). The HCl-salt had m.p. 170-171° (acetone-ether).

10 C<sub>36</sub>H<sub>44</sub>NO<sub>2</sub>CI(558.2)requires: C 77.46 H 7.95 N 2.51 O 5.73 CI 6.35 Found 77.6 7.86 2.42 5.89 6.31

5 I) N-Methyl-N-tert.butyl-3,3-bis-(2,4-dimethoxyphenyl)propylamine (LXXVI), hydrochloride The free base was obtained in 96% yield from the amine (LIX) of Example 6j). The HCl-salt had m.p. 180-190° and seems to be associated with 1/4 mol of water.

C<sub>24</sub>H<sub>36</sub>NO<sub>4</sub>Cl 1/4H<sub>2</sub>O(447.0)requires: C 64.48 H 8.34 N 3.13 O 16.11 Cl 7.93 20 Found 64.5 8.27 3.02 16.2 8.19

m) N-Methyl-N-tert.butyl-3-(2,3-dibenzyloxyphenyl)-3-phenylpropylamine (LXXVII) This was obtained as an oil in 98% yield from the amine (LI) of Example 6b). NMR:  $\delta$  6.9-7.3 (m 18H), 2.1 (s 3H), 1.0 (s 9H).

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n) N-Methyl-N-tert.butyl-3-(2-benzyloxyphenyl)-3-phenylpropylamine (LXXVIII)

This was obtained as an oil in 97% yield from the amine (LII) of Example 6c).

NMR: 6.9-7.3 (m 14H), 5.0 (s 4H), 4.5 (t 1H), 2.2 (s 3H), 0.9 (s 9H).

o) N-Methyl-N-tert.butyl-3-(5-chloro-2-methoxyphenyl)-3-phenylpropylamine
The secondary amine from Example 6o) (25.3 g, 0.076 mol) was refluxed for 18 h with formic acid (9.2 g, 0.2 mol) and 35% formaldehyde solution (8.5 g, 0.1 mol). Work-up gave 25.6 g, (97.5%) crude base. This was dissolved in acetone and treated with an equimolar quantity of oxalic acid in acetone giving beige crystals of the title compound, hydrogen oxalate, m.p. 165°.

C<sub>21</sub>H<sub>28</sub>CINO.C<sub>2</sub>H<sub>2</sub>O<sub>4</sub>(436.0)requires: C 63.37 H 6.94 N 3.21 Cl 8.13 Found 62.7 6.83 3.10 7.97

p) N-(2-Hydroxy-1,1-dimethylethyl)-N-methyl-3-(2-benzyloxyphenyl)-3-phenylpropylamine
This compound was similarly prepared from the compound of Example 5p). It was obtained as a sticky oil which was converted to the free hydroxy compound of Example 9ad).

q) N-1-Adamantyl-N-methyl-3-(2-benzyloxyphenyl)-3-phenylpropylamine This compound was similarly prepared from the compound of Example 5q). It was obtained as a sticky oil which was converted to the free hydroxy compound of Example 9ae) without further purification.

#### Example 8

### Preparation from olefinic precursors

a) N-tert.butyl-3-(2,6-dimethoxyphenyl)-3-hydroxy-3-phenylpropylamine (LXXIX)

A solution of diisopropylamine (10.1 g, 0.1 mol) in dry ether (100 ml) was cooled to -10°. A solution of butyl lithium in hexane (65 ml, 0.1 mol) was added, and the mixture was stirred at -10° for 20 min. A solution of N-ethylidene-tert.butylamine (10 g, 0.1 mol) in dry ether (100 ml) was added and the solution was stirred at 0° for 20 min. After cooling to -30° a solution of 2,6-dimethoxybenzophenone (24.1 g, 0.1 mol) in dry ether (100 ml), containing 30 ml THF, was added. The mixture was then stirred at ambient temperature for 20 h and hydrolized with water. The organic phase was washed with water, dried and evaporated, giving 32 g (94%) of

N-[3-(2,6-dimethoxyphenyl)-3-hydroxy-3-phenylpropylidene]tert.butylamine as an oil. This oil was dissolved in absolute ethanol (250 ml), the solution was cooled to -5°, and NaBH4 (5.7 g, 0.15 mol) was added portionwise. The mixture was stirred at 0° for 1/2 h, then at ambient temperature for 3 h. Most of the solvent was distilled off in vacuum, the residue was treated with water, extracted with ether, washed with water, and extracted with 2N HCl. The extract was washed with ether, basified with NaOH, extracted with ether, dried and evaporated, giving 30 g of the title amine. The HCl-salt had m.p. 203-204° (acetone-ether) and seems to be associated with 1/4 mol of water. 13.52 8.01 3.64 0 65.60 H C21H29NO3.HCl.1/4H2O(384.5)requires: С 13.7 10 8.11 3.64 65.9 Found b) N-tert.Butyl-3-(2,6-dimethoxyphenyl)-3-phenyl-2-propene-1-amine (LXXX) The above amine from step a) (21 g, 0.061 mol) was added to 6.3N H<sub>2</sub>SO<sub>4</sub> (20 ml, 0.126 mol). The mixture 15 was stirred on a boiling water bath for 2 h, cooled, basified, and extracted with ether. The extract was washed, dried and evaporated, giving 17.8 g, (90%) of the title olefin as a clear oil. The HCI-salt had m.p. 220-22°, and was associated with 1/4 mol of water. 9.68 20 0 9.82 CI С 68.82 7.86 Ν 3.82 C<sub>21</sub>H<sub>27</sub>NO<sub>2</sub>.HCl.1/4H<sub>2</sub>O requires: 9.81 7.89 3.92 9.44 68.8 Found c) N-Methyl-N-tert.butyl-3-(2,6-dimethoxyphenyl)-3-phenylpropylamine (LXXXI), hydrogen fumarate 25 The olefinic amine from step b) (16.3 g, 0.05 mol) in methanol (250 ml) containing 0.5 g of a 10% Pd/C catalyst, was hydrogenated at ambient temperature and pressure. The mixture was then filtered through Celaton, the filtrate was taken to dryness, giving 16.3 g (100%) of N-tert.butyl-3-(2,6-dimethoxyphenyl)-3-phenylpropylamine. The HCl-salt had m.p. 244° (ethanol). 30 CI 9.74 3.85 0 8.79 Н 8.31 N C21H29NO2.HCI(363.9)requires: С 69.31 8.29 3.83 9.27 9.75 69.3 Found The above secondary amine, as the free-base, was methylated with formaldehydeformic acid as described in 35 Example 7, giving the tertiary amine in 96% yield. The fumaric acid salt had m.p. 185-190° (acetone). 3.06 20.95 C<sub>26</sub>H<sub>35</sub>NO<sub>6</sub>(457.6)requires: 68.25 H 7.71 3.05 21.6 67.8 7.59 Found 40 Example 9 45 Removal of O-protective groups 50 a) N,N-Diisopropyl-3-(2-hydroxyphenyl)-3-phenylpropylamine (LXXXII), hydrochloride The amine (XLIX) of Example 5k) (20.8 g, 0.064 mol) in methylene chloride (150 ml) was cooled below 0°. A 1N solution of BBr<sub>3</sub> in CH<sub>2</sub>Cl<sub>2</sub> (64 ml, 0.064 mol) was then added dropwise, the solution was then kept in the cooler (5°) for 2-5 days, and volatile material was distilled off at <50°. The residual syrup was basified, extracted with ether, the extract was washed with water, dried and evaporated, giving a viscous syrup. The 55 HCl-salt had m.p. 222° (methanol-ether), yield 31%. 0 4.60 Ci 10.19 72.49 8.69 4.03 C21H29NO.HCI(347.9)requires: С

The following compounds were obtained in the same way.

Found

65

60

5.06

10.3

72.0

8.72

3.74

C <sub>28</sub> H <sub>37</sub> O <sub>5</sub> (467.6)requires: Found	С	71.9 71.8	Н	7.9 8.4		3.00 3.0			1
c) N,N-Diisopropyl-3-(2-hydroxy-5-methylp From the amine (XL) of Example 5c). (	henyl) Crude	-3-phenylp yield 85%	ropy . HC	/lamine ( Cl-salt, m	LXXXIV) p. 209-	, hydrod 210° (ad	chloride etone-	ether	r).
$C_{22}H_{31}NO.HCl.1/4H_2O(366.5)$ requires: Found	С	72.09 72.3	Н	8.95 N 8.95	3.8 3.7		5.46 5.68	CI	
C) N-Methyl-N-tert.butyl-3-(2-hydroxy-5-me From the amine (LXVI) of Example 7b).	thylpl Crud	nenyl)-3-ph le yield 100	enyl 0%.	propylam HCI-salt,	ine (LX m.p. >	XXV), hy 260° (e	droch thanol	loride ).	2
C <sub>21</sub> H <sub>29</sub> NO.HCl(347.4)requires: Found	С	72.49 72.7	Н	8.69 8.59		4.00 3.8			10
e; N,N-Diisopropyl-3,3-bis-(2-hydroxyphen From the amine (XXXVIII) of Example 5	yl)pro ia). Ci	pylamine (I rude yield (	LXX 57%	XVI), hyd o. HCI-sal	rochlori t, m.p.	<u>de</u> 257° (et	hanol-	ether	).
C <sub>21</sub> H <sub>29</sub> NO <sub>2</sub> .HCl(363.9)requires: Found	С	69.31 (69.3	Н	8.31 N 8.37	3.8 3.9		8.79 9.23	CI	
f) N-Methyl-N-tert.butyl-3,3-bis-(2-hydroxy From the amine (LXVII) of Example 7c	pheny ). Cru	l)propylam de yield 10	ine 10%,	(LXXXVII) m.p. 19	, hydro 0°. HCI	chloride -salt, m.	p. <b>252</b> °	eth (eth	nar
C <sub>20</sub> H <sub>27</sub> NO <sub>2</sub> .HCl(349.9)requires: Found	С	68.65 68.4	Н	8.0 8.0		4.0 4.1			1
gj N.N-Diisopropyl-3-(2-hydroxy-4-methylp From the amine (XLI) of Example 5d).	ohenyl Crude	)-3-phenylp yield 90%	orop 6. H	ylamine ( Cl-salt, m	LXXXVI 1.p. 217	II), hydro ° (ethan	ochlori	de	
C <sub>22</sub> H <sub>31</sub> NO.HCl.1/4H <sub>2</sub> O(366.5)requires: Found	С	72.09 72.3	Н	8.96 I 8.91	3.8 3.9		5.46 5.27	CI	
						.1 .1.1			
hj N.N-Diisopropyl-3,3-bis-(2-hydroxy-5-m From the amine (XLII) of Example 5e).	Crud	e yield 93%	ylan Ио, п	n.p. 166°	HCl-sa	ult, m.p.	220° (	ethar	nol
C <sub>22</sub> H <sub>33</sub> NO <sub>2</sub> .HCl(392.0)requires: Found	С	70.47 70.6	Н	8.7 8.7		3.5 3.7			8
	rude '	thylphenyl) yield 79%, i	m.p.	199-201°	(XC), n	ydrochic HCl-salt,	m.p. 2	20° (a	ace
N-Methyl-N-tert.butyl-3,3-bis-(2-hydroxy From the amine (LXIX) of Example 7e). C									

From the amine (LXVIII) of Example 7d). Crude yield 100%. HCI-salt, m.p. 240° (ethanol).												
C <sub>21</sub> H <sub>29</sub> NO.HCl(347.9)requires: Found	С	72.49 72.5	н	8.69 8.75	N	4.03 4.06	0	4.60 4.90	CI	10.19 10.1	5	
	\ <b></b>		l\n=on	ulomina	. /۷0	an byd	rochl	orida				
k) N,N-Diisopropyl-3-(4-fluorophenyl)-3-(2 From the amine (XLVII) of Example 5j	. Crud	de yield 7	/2%.	HCI-sal	t, m.	o. 183°	(acei	one-eth	nanol	).	10	
C <sub>21</sub> H <sub>27</sub> FNO.HCl(364.9)requires: C Found		69.12 69.1	Н			7.73 8.09	N			3.83 3.82		
I) N,N-Diisopropyl-3-(2,4-dihydroxyphenyl From the amine (XLV) of Example 5h).	)-3-ph Crude	enylpropy yield 31%	/lamin /o. HC	e (XCII I-salt, n	I), hy n.p. 2	drochic 05-210	oride (eth	anol-ac	etone	e-ether).	15	
C <sub>21</sub> H <sub>29</sub> NO <sub>2</sub> .HCl(363.9)requires: Found	С	69.31 69.5	Н	8.31 8.33	N	3.85 3.72	0	8.79 8.91	CI	9.74 9.87	20	
m) N-(1,1-Dimethylpropyl)-N-methyl-3,3-b From the amine (LXXII) of Example (ethanol-acetone-ether).	is-(2-h 7h). C	ydroxy-5 rude yiel	-meth d 100	ylpheny )%, m.	/l)pro p. 19	pylamir 90-195°	ie (XC . HCl	CIV), hy -salt, m	droci	nloride 235-240°	25	
$C_{23}H_{33}NO_2$ .HCI(392.0)requires: Found	С	70.47 70.0	Н	8.74 8.96	N	3.57 3.54	0	8.16 8.11	CI	9.05 9.19	30	
n) N-Methyl-N-tert.butyl-3-(2,4-dihydroxy) From the amine (LXXIII) of Example 7i).	ohenyl Crude	)-3-pheny yield 78%	iprop , m.p.	ylamine 260°. ł	(XC IBr-s	V), hydalt, m.p.	robro . > 26	mide 0° (etha	anol).		35	
C <sub>20</sub> H <sub>25</sub> NO <sub>2</sub> .HBr(394.4)requires: Found	С	60.9 60.8	Н	7.16 7.18	N	3.55 3.29	0	8.11 8.38	Br	20.27 20.2		
o) N,N-Diisopropyl-3,3-bis-(2,4-dihydroxy From the amine (XLVI) of Example 5i). T a satisfactory elemental analysis becaus	he HC	I-salt, con	sistin	g of an	amor	ochloric phous t	l <u>e</u> orown	powde	r, did	not give	40	
p) N-Methyl-N-tert.butyl-3,3-bis-(2,4-dihyl From the amine (LXXVI) of Example 7I). elemental analysis because of incomplet	Crude	yield 870	pylan ⁄o, m.;	nine (X o. 260°.	CVII), The	hydrod HCI-sal	hlorid did r	<u>de</u> iot give	a sat	isfactory	45	
q) N,N-Diisopropyl-3-(2,5-dihydroxypheny The amine (XLIII) of Example 5f) in the containing 5 g of a 5% Pd/C catalyst was reaction was complete. The mixture was finacetone and treated with ethereal HCl, gimethanol gave white crystals, m.p. 260°	ne fori hydro Itered, ving 1	n of the ogenated the filtrat	free I at am e was	base (3 bient te taken t	32 g, empe to dry	0.063 r rature a ness, th	nol) i ind pr ie res	n meth: essure. idue wa	Afte s dis:	r 2 h the solved in	50	
C <sub>21</sub> H <sub>29</sub> NO <sub>2</sub> .HCl.1/4H <sub>2</sub> O(368.6)requires: Found	C	68.44 68.4	Н	8.36 8.40	N	3.80 3.60	0	9.77 10.3	CI	9.62 9.42	55	
The following compounds were prepa											60	
r) N-Methyl-N-tert.butyl-3-(2,5-dihydroxyg From the amine (LXXIV) of Example 7	henyl j). Cr	)-3-pheny ude yield	90%.	ylamine HCl-sa	(XC	X), hyd .p. >2	rochl 70° (r	oride nethano	ol-wa	ter).		
											65	

	C <sub>20</sub> H <sub>27</sub> NO <sub>2</sub> .HCI(349.9) requires: Found	С	68.65 68.9	Н	8.06 8.02	N	4.00 3.93	0	9.14 9.60	CI	10.13 10.5
5	s) N.N-Diisopropyl-3,3-bis-(2-hydroxy-4-me From the amine (XLIV) of Example 5g).	thylph Crud	nenyl)pro e yield 1	pylan 00%.	nine (C HCl-sa	), hyc alt, m	rochlor p. 253°	ide ' (me	thanol-	ether	).
10	C <sub>23</sub> H <sub>33</sub> NO <sub>2</sub> .HCI(392.0) requires: Found	С	70.47 70.5	Н	8.74 8.74	N	3.57 3.55	0	8.16 8.47	CI	9.05 8.03
15	t) N-Methyl-N-tert.butyl-3,3-bis-(2-hydroxy- From the amine (LXXV) of Example (methanol-acetone).	-4-met 7k). (	thylphen Crude yi	yl)pro eld 9	pylamir 7%, a	ne (Cl yello	), hydro w pow	ochlo der.	ride HCl-sal	lt, m.	p. 260°
20	C <sub>22</sub> H <sub>31</sub> NO <sub>2</sub> .HCl(378.0)requires: Found	С	69.91 69.9	Н	8.54 8.68	N	3.71 3.67	0	8.47 8.85	CI	9.38 9.24
.25	u) N,N-Diisopropyl-3-(2,3-dihydroxyphenyl) From the amine (XXXIX) of Example 5b	-3-phe ). Cru	enylprop ide yield	ylamir 100%	ne (CII) 6. HCI-	, hydi salt, i	n.p. 17	<u>de</u> 4-176	° (acet	one).	
	C <sub>21</sub> H <sub>29</sub> NO <sub>2</sub> .HCl(363.9)requires: Found	С	69.31 69.5	Н	8.31 8.33	N	3.85 3.66	0	8.79 9.37	ÇI	9.74 9.63
30	w) N-Methyl-N-tert.butyl-3-(2,3-dihydroxyp) From the amine (LXXVII) of Example 7m) heating, (methanol-acetone).	henyl) . Cruc	-3-pheny le yield 1	/lprop 00%,	ylamine a white	e (CIII	), hydro der. HC	ochlo I-salt	<u>ride</u> , m.p. 2	09-21	0°, slow
35	C <sub>20</sub> H <sub>27</sub> NO <sub>2</sub> .HCl.1/4H <sub>2</sub> O(358.9)requires: Found	С	66.92 66.9	Н	8.14 8.12	N	3.90 3.76	0	11.14 11.8	CI	9.88 9.74
40	x) N-methyl-N-tert.butyl-3-(2-hydroxyphen From the amine (LXXVIII) of Example 7	yl)-3-p n). Cı	henylpro rude yiel	pylan d 100	nine (C %. HC	IV), h I-salt,	ydrochl m.p. 2	oride 55° (	acetone	e-ethe	er).
.15	C <sub>26</sub> H <sub>27</sub> NO.HCl(333.9)requires: Found	С	71.9 71.9			8.45 8.43	N	4.2 4.0			10.62 10.5
50	y: N-Methyl-N-tert.butyl-3-(2,6-dihydroxypl From the amine (LXXXI) of Example 8d	henyl) c) with	-3-pheny n BBr <sub>3</sub> , i	Iprop	ylamine yield.	(CV)	, hydro alt, m.p	chlor . 170	<u>ide</u> ° (etha	nol-e	ther).
au	C <sub>29</sub> H <sub>27</sub> NO <sub>2</sub> .HCl.1/2H <sub>2</sub> O(358.9)requires: Found	С	66.93 67.4	Н	8.14 8.28	N	3.40 3.63	0	11.14 10.9	CI	9.87 9.99
55	zj N.N-Diisopropyl-3-(5-chloro-2-hydroxyp The base from Example 5m) (11.7 g, 0.03	2 mol)	was trea	ited w	ith pyric	dine (1	7.6 g, 0.	096 m	nol) and	conc	. HCI (13
60	g). The mixture was taken to dryness in vac h. The melt was cooled somewhat, water cooled. 2 N HCl was added, the salt was fill salt m.p. 200°. Recrystallization from acet	was a tered	dded, th off, wash	e mix ed wi	ture wa th 2 N F	is dig ICI an	ested in d dried	n a bo , givin	oiling w 11.0 q	ater t g (90°	oath and /o) white
65	C <sub>21</sub> H <sub>28</sub> CINO.HCI(382.4)requires: Found	С	65.9 66.0			7.64 7.88	N		.66 .63	CI	18.54 18.3

aa) N-Methyl-N-tert.butyl-3-(5-chloro-2-hydroxyphenyl)-3-phenylpropylamine  The free base from Example 7o) (10.5 g, 0.03 mol) was treated with pyridine (7.0 g, 0.09 mol) and conc. HCl (12 g). The mixture was taken to dryness in vacuum and the residue was heated in an oil-bath at 205-215° for 1 1/2 h. The melt was cooled somewhat, excess of 2 N NaOH was added, the mixture was extracted with ether, the extract was washed with water, dried and evaporated giving 7.5 g (88%) crude syrup. This was dissolved in ether and treated with ethereal HCl giving 8 g (83%) of hydrochloride salt. Recrystallization from acetone-2 N HCl gave the hydrochloride of the title compound, m.p. 260°.														
C <sub>20</sub> H <sub>26</sub> CINO.HCl(368.4)requires: Found	С	65.21 65.0	Н		7.39 7.30	N	3.80 3.73			19.25 18.9	10			
ab) N-[3-(2-Hydroxyphenyl)-3-phenylpropyl]-2,2,5,5-tetramethylpyrrolidine  The crude amine from Example 5n) was hydrogenolysed as described in Example 9q). The free amine was obtained as an oil which was converted to the hydrochloride and crystallized from 2-propanol. M.p. 250°C.  C23H31NO.HCI(374.0)requires:  C 73.86 H 8.63 N 3.75 O 4.28 Cl 9.48														
C23H31NO.HCI(374.0)requires:       C       73.86       H       8.63       N       3.75       O       4.28       CI       9.48         Found       73.8       8.71       3.59       4.80       9.45														
ac) N-[3-(2-Hydroxyphenyl)-3-phenylpropyl The benzyloxy compound from Example base was converted to the hydrochloride s melts with decomposition at about 150°C	50) v emihy	vas hydro	ogen	olysed	as de	scribed	in Ex	ample one. Ti	9q). пе со	The free mpound	25			
$C_{24}H_{33}NO_2.HCl.1/2H_2O(413.0)$ requires: Found:	С	69.79 70.0	н	8.54 8.67		3.39 3.47	0	9.68 9.98	CI	8.58 8.13	30			
ad) N-(2-Hydroxy-1,1-dimethylethyl)-N-met The benzyloxy compound from Example obtained as a glassy mass, was converted t precipitation from ethanol with ether.	7p) w	as hydrod	geno	ysed a	as des	cribed ir	า Exar	nple 90 amorp	զ). Th ohous	e amine, solid on	35			
$C_{20}H_{27}NO_2$ .HCI(349.9)requires: Found:	С.	68.65 68.25	Н	8.06 8.18		4.00 3.98	0	9.15 9.12	CI	10.13 10.0	40			
ae) N-1-Adamantyl-N-methyl-3-(2-hydroxy) The benzyloxy compound from Example hydroxyamine was obtained as a glassy ma of hydrogen chloride in ether. The hydrochlo	7q) ss. lt	was hydr was disso	ogen olved	olysed in anh	i as de ydrou:	s ether a	and tre	ated v	vith a	n excess	45			
C <sub>26</sub> H <sub>33</sub> NO.HCl(412.0)requires: Found:	С	75.79 75.3	Н	8.32 8.01		3.40 3.22	0	3.88 3.45	CI	8.61 8.96	50			
		Example	<u>10</u>								55			
	Red	uction of	amic	<u>les</u>							60			
a) N,N-Diisopropyl-3-(2-methoxy-5-methyl 3-(2-Methoxy-5-methylphenyl)-3-phenyl Chem. Soc. 1956 1382) and thionyl chlorid chloride is distilled off under reduced pre	oropio e (50	nic acid ml) are h	(12.8 eatec	g, 0.0 I on a	05 moi water	bath for	3 h. T	he exc	cess o	of thionyl	65			

nyipropionyl chloride is dissolved in 50 ml of dichloromethane and added dropwise to a stirred solution of diisopropylamine (20.2 g, 0.20 mol) in 200 ml of dichloromethane at about 0°C. The solution is left for 2 h, the solvent is distilled off and the remaining material is treated with water. The solid product consisting of N,N-diisopropyl-3-(2-methoxy-5-methylphenyl)-3-phenylpropionamide is filtered off, dried and added in small portions to a stirred suspension of lithium aluminium hydride (6.0 g, 0.16 mol) in dry ether (700 ml). The mixture is refluxed for 2 days. Excess of hydride is destroyed by the careful addition of water, the ether layer is separated and dried with anhydrous sodium sulfate. After filtration the solution is added to a solution of excess fumaric acid in ether. The precipitated salt is collected and crystallized from 2-propanol. The hydrogen fumarate melts at 176°C.

b) N-Methyl-N-tert.butyl-3-(2-methoxy-5-methylphenyl)-3-phenylpropylamine was similarly prepared. The hydrochloride melts at 161°C.

Example 11

a) N-Methyl-N-tert.butyl-3-(5-chloro-2-hydroxyphenyl)-3-phenylpropylamine

A solution of chlorine (7,1 g, 0.10 mol) in acetic acid (500 ml) is added dropwise to a stirred solution of N-methyl-N-tert.butyl-3-(2-hydroxyphenyl)-3-phenylpropylamine (29.7 g, 0.10 mol) in acetic acid (200 ml) with stirring. After 2 h the solvent is distilled off under reduced pressure and the crude hydrochloride left is recrystallized from 2-propanol. Melting point 260°C.

b) N,N-Diisopropyl-3-(5-chloro-2-hydroxyphenyl)-3-phenylpropylamine is similarly prepared. The hydrochloride melts at 202-3°C.

#### Example 12

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### Separation of (+)- and (-)-enantiomers

( $\pm$ )-N,N-Diisopropyl-3-(2-hydroxyphenyl)-3-phenylpropylamine (31.1 g, 0.10 mol) is dissolved in 300 ml of ethanol. A solution of L(+)-tartaric acid (15.0 g, 0.10 mol) in 400 ml of ethanol is added. The mixture is heated a few minutes in a boiling water bath and seeded with crystals obtained by cooling and scratching a small sample of the main solution. The mixture is chilled at about 4°C over-night whereupon the crystalline precipitate is filtered off, washed with cold ethanol and recrystallized repeatedly from ethanol. The pure (-)-N.N-diisopropyl-3-(2-hydroxyphenyl)-3-phenylpropylamine hydrogen L-(+)-tartrate thus obtained has  $[\alpha]_0^{20}$   $\frac{10.6^{\circ}}{10.6^{\circ}}$  (c = 5% in methanol). The free amine is obtained by alkalisation of an aqueous solution, extraction into ether, drying and evaporation of the solvent. Sticky oil,  $[\alpha]_0^{20}$  -5.4° (c = 5% in methanol).

(+)-N,N-Diisopropyl-3-(2-hydroxyphenyl)-3-phenylpropylamine is similarly prepared using D-(-)-tartaric acid. The hydrogen-D-(-)tartrate has  $[\alpha]_{\rm D}^{20}$  + 10.0°. The free amine has  $[\alpha]_{\rm D}^{20}$  + 5.6°, both measured as 5% solutions in methanol.

Example 13 (continuation of Example 1)

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### Preparation of 4-phenyl-3,4-dihydrocoumarins

g) 4-(2-Methoxyphenyl)6-methyl-3,4-dihydrocoumarin (CVI)

A mixture of 2-methoxycinnamic acid (178 g, 1.0 mol), p-cresol (108 g, 1.0 mol), and p-toluenesulphonic acid monohydrate (47.5 g, 0.25 mol) was stirred on a boiling water-bath for about 2 h during which time the system was evacuated with a waterpump to remove formed water. The solid was then broken up and washed copiously with water. The granular material was then stirred with a large volume of saturated NaHCO<sub>3</sub> solution containing some 10% acetone. The product was filtered off, washed, dried and recrystallised from acetone affording 167 g (62,5%) white crystals of the desired lactone, m.p. 140°.

C <sub>16</sub> H <sub>13</sub> O <sub>3</sub> (288.7) requires:	С	66.56	Н	4.54	0	16.62
Found:		66.8		4.45		16.5

h) 6-Chloro-4-(2-methoxyphenyl)-3,4-dihydrocoumarin (CVII) was prepared in a similar way in 49% yield from 2-methoxycinnamic acid and p-chlorophenol, the reaction temperature being 130° in this case. M.p. 172-173° (acetone).	_
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EMI ID = 34/2 HE = 15 WI = 125 TI = TAB	10
Example 14 (continuation of Example 2)	15
Example 14 (continuation of Example 2)	
Preparation of 3,3-diphenylpropionic acid esters	20
I) Methyl-3-(2-methoxyphenyl)-3-(2-methoxy-5-methylphenyl)propionate (CVIII) was obtained as an oil in 75% yield from the lactone CVI of Example 13g in the manner described for the ester VI of Example 2a).	
m) Methyl-3-(5-chloro-2-methoxyphenyl)-3-(2-methoxyphenyl)propionate (CIX) was obtained as an oil in the same way in 97% yield from the lactone CVII of Example 13.	25
Example 15 (continuation of Example 3)	30
Preparation of 3,3-diphenylpropanols	35
m) 3-(5-Chloro-2-methoxyphenyl)-3-(2-methoxyphenyl)propanol (CX) was obtained in 84% yield from the ester CIX of Example 14m in the manner described for the propanol XVI of Example 3a), except that the reduction was carried out in toluene with a 10% molar excess of a 3.4 M toluenic solution of sodium bis(2-methoxyethoxy)aluminium hydride (SMEAH) instead of LiAlH4. M.p. 70-72° (IPE).	40
n) 3-(2-Methoxyphenyl)-3-(2-methoxy-5-methylphenyl)propanol (CXI) was obtained in the same way in quantitive yield from the ester CVIII of Example 14I). The product consisted of a golden oil of 89% purity according to GC.	45
Example 16 (continuation of Example 4)	50
Preparation of 3,3-diphenylpropyl-p-toluenesulphonates	
n) 3-(2-Methoxyphenyl)-3-(2-methoxy-5-methylphenyl)propyl-p-toluenesulphonate (CXII) was prepared in the same way as the tosylate XXVII of Example 4a) in quantitative yield from the propanol CXI of Example 15n) using CH <sub>2</sub> Cl <sub>2</sub> as solvent instead of chloroform. M.p. 101° (ether/IPE).	55
C <sub>25</sub> H <sub>28</sub> O <sub>5</sub> S (440.57) requires:       C       68.16       H       6.41       S       7.28         Found:       68.3       6.51       7.20	60
o) 3-(5-Chloro-2-methoxyphenyl)-3-(2-methoxyphenyl)propyl-p-toluenesulphonate (CXIII) was obtained in the same way in quantitative yield from the propanol CX of Example 15m. M.p. 97-98°	65

(acetone/IPE).

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C24H25CIO5S (460.92)requires:	С	62.54 H	5.47 S	6.94 CI	7.69
Found:		63.0	5.65	6.95	7.70

Example 17 (continuation of Example 5)

# Preparation of tertiary 3,3-diphenylpropylamines

r) N,N-Diisopropyl-3-(5-chloro-2-methoxyphenyl)-3-(2-methoxyphenyl)propylamine (CXIV) was obtained as an oil in 94% yield from the tosylate CXIII of Example 16o) in the manner described for the amine XXXVIII of Example 5a). Purity by GC = 99.9%.

s) N,N-Diisopropyl-3-(2-methoxyphenyl)-3-(2-methoxy-5-methylphenyl)propylamine (CXV) was obtained in the same way in 49% crude yield from the tosylate CXV of Example 16n). After chromatographic purification on an Si-gel 60 column (eluation with light petroleum), the product (oil) had a purity of 100% according to GC.

t) N-[(2-Benzyloxy-5-methyl)-3-phenyl]-2,2,5,5-tetramethylpyrrolidine (CXVI) was prepared from 3-(2-benzyloxy-5-methyl)-3-phenylpropyl tosylate and 2,2,5,5-tetramethylpyrrolidine following the directions given in Example 5a). It was obtained as a sticky oil which was converted to the free hydroxy compound of Example 20aj).

### Example 18 (continuation of Example 6)

# Preparation of secondary 3,3-diphenylpropylamines

p) N-tert.Butyl-3-(5-chloro-2-methoxyphenyl)-3-(2-methoxyphenyl)propylamine (CXVII) was prepared in quantitative yield from the tosylate CXIII of Example 16o) in the manner described for the amine L of Example 6a). The HCI-salt had m.p. > 260°.

C <sub>21</sub> H <sub>28</sub> CINO <sub>2</sub> .HCI (398.38) requires:	С	63.3	Н	7.34	N	3.52 C	17.80
Found:		63.2		7.46		3.49	17.4

g) N-tert.Butyl-3-(2-methoxyphenyl)-3-(2-methoxy-5-methylphenyl)propylamine (CXVIII) was obtained in a similar way in 89% crude yield from the tosylate CXII of Example 16n). The HCl-salt had m.p. 225°.

C <sub>22</sub> H <sub>31</sub> O <sub>2</sub> N.HCl (377.97)										
Requires:	С	69.91	Н	8.54	Ν	3.71	Cl	9.38	0	8.47
Found:		69.8		8.73		3.60		9.45		8.79

Example 19(continuation of Example 7)

Preparation of tertiary 3,3-diphenylpropylamines from secondary amines

r) N-Methyl-N-tert.butyl-3-(5-chloro-2-metwas prepared in 89% yield from the amine	hoxyph	enyl)-3-(2-me	thoxypheny	)propyl	amine (CXI)	<u>()</u> ne amine l	LXI of
Example 7a). The HCI-salt was prepared	by trea	iting an aceto	nic solution	of the	free base v	ith contra	acted
hydrochloric acid. M.p. 130°.	-						5
0 11 010 1110111 0 (400 40)							3
C <sub>22</sub> H <sub>30</sub> ClO <sub>2</sub> N.HCl.H <sub>2</sub> O (430.42) Requires:	С	61.39 H	7.74	N	3.25 C	I 16	3.47
Found:	Ū	62.0	7.93		3.26	16	.5
, cana.							10
							70
s) N-Methyl-N-tert.butyl-3-(2-methoxyphe	nyl)-3-	2-methoxy-5	methylphen	yl)propy	lamine (CX	<u>X)</u>	
was prepared in a similar way in 98% yiel	d from	the amine CX	VIII of Exam	ple 18q	). The free t	oase (oil) l	had a
purity of 96% by GC.							15
_		., ., .	<b>5</b>				
Example	20 (c	ontinuation of	Example 9	)			
							20
Ren	noval o	f O-protective	groups				
af) N,N-Diisopropyl-3-(2-hydroxyphenyl)-	3-(2 <b>-</b> hyc	droxy-5-methy	iphenyl)proj	oylamine	(CXXI)		<i>25</i>
The amine CXV from Example 17s) (2)	6.5 a. C	).072 mol) in 1	methanol wa	as treat	ed with a s	light exce	ss of
concentrated hydrochloric acid. The mixture mol) was added and the mixture was then	re was	taken to dryne	ess in vacuu or 1 1 h The	m, pyria • mixtur	inium chiori e was coole	de (25.4 g d to abou	, 0.22 t 80°.
acetone (20 d) was added followed by add	lition of	little water. Ti	ne salt was f	iltered c	off, washed \	vith dilute	a HCI
and dried. Recrystallisation from absolut	e ethar	ol/ether gave	17.5 g (64.	3%) of	a white sal	t, m.p. >	250°. <i>30</i>
Purity by $GC = 100\%$ .							
0 H NO HOL(277.07)							
C <sub>22</sub> H <sub>31</sub> NO <sub>2</sub> .HCl (377.97) Requires:	С	69.91 H	8.54 N	3.71	O 8.47	CI S	9.38
Found:		69.8	8.65	3.57	8.76	5 9	9.51 <sup>35</sup>
ag) N,N-Diisopropyl-3-(5-chloro-2-hydrox	ypheny	i)-3-(2-hydrox	yphenyl)pro	pylamin	e (CXXII		0149 40
was prepared in the same way in 37% yie	ld from	the amine CX	IV of Examp	ie 17r).	The HCI-sai	t nad m.p.	. 214° 40
(ethanol).							
C <sub>21</sub> H <sub>28</sub> NO <sub>2</sub> .HCl (398.38)				•			
Requires:	С	63.31 H	7.34 N	3.52	O 8.03		7.80
Found:		63.1	7.34	3.40	8.1	5 1	7.8 <sup>45</sup>
	n .	(O. b l	41l l		damina (CV	VIII\	
ah) N-methyl-N-tert.butyl-3-(2-hydroxyphwas prepared in the same way in 30% yis	enyi)-3	the amine C	X of Examp	yi)propy le 19s).	The HCI-sal	t had m.p.	. 240° <i>50</i>
(acetone).	ila non	1110 0111110 01		,.		•	
(400000)							
C <sub>21</sub> H <sub>29</sub> NO <sub>2</sub> .HCl (363.94) requires:	С	69.3 ⊦		N			7.74
Found:		69.0	8.35		3.65	9	).76 - <i>55</i>
		h () () () ()	ت جا جرين مسامر	d\	omine (CV)	(1) (1)	
ai) N-Methyl-N-tert.butyl-3-(5-chloro-2-h was prepared in the same way in 24%	droxyr	onenyi)-3-(2-h)	CXIX of F	npropyl	19r), M.D.	> 250°.	
was prepared in the Same way in 2490	, ICIU II	on the annie					60
C <sub>20</sub> H <sub>26</sub> CINO <sub>2</sub> .HCI (384.36) requires:	С	62.50 H	7.08	N	3.65	CI 18	8.45
Found:		62.5	7.09		3.63	18	B.4

aj) N-[3-(2-Hydroxy-5-methylphenyl)-3-phenylpropyl]-2,2,5,5-tetramethylpyrrolidine (CXXV) was obtained when the O-benzylated amine CXVI of Example 17t) was hydrogenolyzed as described in Example 9q. The hydrochloride melts at 240°. 74.29 8.83 3.61 CI 19.14 C24H34CINO (388.0) requires: 73.9 8.90 3.52 9.48 Found: 10 Example 21 (continuation of Example 10) Reduction of amides 15 N,N-Diisopropyl-3-(2-methoxyphenyl)-3-phenylpropionamine N,N-Diisopropyl-3-(2-methoxyphenyl)-3-phenylpropionamide was obtained as o pale yellow oil in quantitative yield from 3-(2-methoxyphenyl)-3-phenylpropionic acid in the manner described for the amide of Example 10a). This amide (27 g, 0.08 mol) in toluene (50 g) was added dropwise under r.t. to a 3.4 M toluenic solution of SMEAH (50 g, 0,17 mol) diluted with an equal weight of toluene. The mixture was stirred at 60-70° for 2 h, cooled, treated with excess od 2N NaOH. The organic phase was separated, washed with water and extracted with 2N HCl. The acidic extract was washed with ether, basified, extracted with ether, dried and evaporated giving 17.1 g (66%) free base. This was dissolved in acetone (75 ml) and treated with 6.6 g fumaric acid 25 dissolved in methanol, affording 20 g of the fumaric acid salt, m.p. 163-164°. C<sub>22</sub>H<sub>31</sub>ON.C<sub>4</sub>H<sub>4</sub>O<sub>4</sub> (441.58) requires: 70.72 H 7.99 3.17 Ω 18.12 70.7 7.96 3.13 18.0 Found: 30 Example 22 35 Separation of (+)- and (-)-enantiomers 40  $(\div)\text{-N.N-Diisopropyl-3-} (2\text{-hydroxy-5-methylphenyl})\text{--3-phenylpropylamine hydrogen } tartrate$ The racemic amine (LXXXVIII of Example 9g) (48.8 g, 0.15 mol) was dissolved in 500 ml of 95% ethanol and mixed with a solution of L(+)-tartaric acid (22.5 g, 0.15 mol) in 500 ml of ethanol. The mixture was left over night at +4°. The precipitated salt was collected by filtration and washed with ethanol and ether. The yield of +29.5° (C 5%, methanol) was 34,3 g. Two recrystallisations from ethanol afforded 45 crude salt with [α] 😤 🕫 21.8 g with [α] 📆 🕫 +36.0°. C: 65 66 H 7.84 N 2.95 0 23.55 C26H37NO7 requires: Found: 65.9 8.06 2.90 23.5 50 !-)-N,N-Diisopropyl-3-(2-hydroxy-5-methylphenyl)-3-phenylpropylamine hydrogen D(-)-tartrate was similarly prepared using D(-)-tartaric acid. [α] 356 С 65.6 8.00 2.83 55 Н 23.6 Found: Several of the compounds according to the invention were tested with regard to anti-cholinergic, anti-noradrenaline, and anti-calcium effects, toxicity and effect on the heart rate. The test procedures are described below, and the test results are reported in Table 1. For comparison purposes the testing also included the commercially available drug terodiline and a structurally similar compound, N,N-dimethyl-

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activity and the undesired side-effects.

3-(2-methoxyphenyl)-3-phenylpropylamine, disclosed as an antidepressant in US-A-3.446.901, GB-A-1.169.944, and GB-A-1.169.945. The test results clearly show that the compounds according to the invention are superior to the known compounds especially as regards selectivity between the desired anti-cholinergic

a) Anticholinergic activity on isolated urinary bladder

Male guinea-pigs, weighing 250-350 g, were killed by a blow on the head and exsanguinated. The urinary bladders were quickly removed and placed in Na<sup>+</sup>-Krebs, in which they were kept throughout the dissection procedure. The bladders were dissected free from adherent fat and connective tissue before they were cut open by an incision on each side from the base towards apex. The mucosa was carefully removed with a pair of scissors. Four strips, approximately 3-5 mm long were prepared by cutting in a parallel direction to the longitudinal muscle fibres, on each half of the bladder.

The bladder strips were immediately mounted vertically in 5 ml organ baths containing Na\*-Krebs solution aerated with carbogene gas to maintain the pH at about 7.4. The temperature, 37°C, was thermostatically controlled by a Lauda MS3 thermostatic circulator. The preparations were suspended between two hooks, one of which was connected to a Grass Instruments FTO3 force transducer. The isomeric tension of the preparations was recorded by a Grass polygraph model 79D. The resting tension was applied to approximately 5 mN. The strips were allowed to stabilize for at least 45 minutes. During this period the resting tension was adjusted to 5 mN and the preparations were repeatedly washed.

In the preliminary experiments concentration - effect curves for carbachol (carbamylcholin chloride) were studied, in order to determine a suitable agonist concentration for inhibition studies with antagonist. The carbachol concentration chosen, 3x10<sup>-6</sup>M, produced a submaximal contractant response (70%). In the inhibition studies, the strips were contracted with carbachol (3x10<sup>-6</sup>M) every 15 minutes. The strips were washed three times after every agonist addition. This procedure was repeated until a reproducible contractant response was observed. A variation of about 10% for three subsequent contractions was accepted as reproducible.

Initially each antagonist was tested in a concentration of 10<sup>-6</sup>M, on two bladder-strips from different guinea-pigs. When a reproducible response with 3x10<sup>-6</sup>M carbachol was obtained, the strips were incubated with the antagonist for 15 minutes before the next carbachol was added. If the antagonist produced more than 50% inhibition of the response to carbachol, a complete concentration-inhibition curve was also made. In the complete inhibition curves, the strips were then incubated for 60 minutes with a fixed concentration of the antagonist before the next addition of carbachol. The effect of the antagonists was calculated as per cent inhibition of the mean of the initial agonist-induced contractions. To generate concentration-inhibition curves the antagonists were studied in 6-8 concentrations and for each concentration a fresh preparation was used, i.e. the strips were only exposed to the antagonist once before they were discarded.

# b) Antagonistic effect to noradrenaline and calcium on the portal vein

### Preparation of isolated portal vein from rat

Animals: Albino, male rats, weighing about 200 g.

Bath volume: 5 ml

Buffer: Na+-Krebs, modified by K.E. Andersson

Temperature: 37°C

Gas: Carbogene (93.5% O<sub>2</sub> + 6.5% CO<sub>2</sub>)

Muscle tension: 0.5 g

The rat is killed by a blow on the neck and decapitated. The abdomen is opened, the vein is dissected free from fat, cut open longitudinally and mounted in an organ bath. Changes in isometric tension is registered by a force displacement transducer, connected to an amplifier and a writing oscillograph.

### Noradrenaline - antagonism on portal vein

Doses: Noradrenaline 3x10<sup>-7</sup> M

The chosen doses give about 70% of maximal response. The agonist is added to the bath at 10-minutes intervals. When reproducible contractions are obtained a fixed concentration of the test substance is added to the bath. After an incubation period of 10 minutes noradrenaline is added. The next concentration of the test substance is added when the original response of the agonist is obtained.

The antagonistic effect of the substance is calculated as per cent inhibition of the mean response by three preceding doses of the agonist.

### Ca - antagonistic effect on portal vein

10 mM K\*-solution is added to the Krebs buffer to stabilize the spontaneous myogenic activity of the vein. The amplitude of the muscle contractions is measued. The test substance is added to the bath in cumulative doses until total inhibition is obtained.

#### c) Histamine - antagonism on isolated ileum

Preparation of isolated ileum from guinea pigs

Animals: Guinea pigs of both sexes, weighing about 350 g.

Bath volume: 5 ml

Buffer: Na\*-Krebs, modified by K.E. Andersson

Temperature: 37°C

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Gas: Carbogene (93.5% O<sub>2</sub> + 6.5% CO<sub>2</sub>)

Muscle tension: 0.5 g

The guinea pig is killed by a blow on the neck and decapitated. The abdomen is opened and about 2 cm of the ileum is cut off about 15 cm above the ileocaecal junction. The piece of ileum is washed with buffer and mounted in an organ bath. Changes in isometric tension is recorded by a force displacement transducer, connected to an amplifier and a writing oscillograph. Dose:  $5x10^{-7}$  M of histamine

The chosen dose of histamine gives about 70% of maximal response. The agonist is added to the bath at 3-minutes intervals. When reproducible contractions are obtained a fixed concentration of the test substance is added to the bath. After an incubation period of 2-10 minutes a new contraction is induced by histamine. The next concentration of the test substance is added when the original response of the agonist is obtained.

The agonistic effect of the test substance is calculated as per cent inhibition of the mean response by three preceding doses of histamine.

d) Acute toxicity in mice

The antagonists to be tested were dissolved in 0.9% NaCl. If they were not soluble in 0.9% NaCl they were dissolved in double distilled water. The solutions were prepared on the day of the experiment.

### Procedure

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White male mice, 25 g, were placed in a mouse holder. The tested compounds were given as i.v. bolus doses in one of the four tail-veins, with a volume of 0.01 ml/g mouse. Each substance concentration was given to a group of four mice. 4-5 different concentrations of the antagonists were made and tested.

The acute lethal dose (LD<sub>11</sub>) was the lowest concentration of the anticholinergic drug where 4 mice of 4 tested died within 5 minutes after an i.v. bolus dose.

 $LD_{50}$ -interval: The  $LD_{50}$ -interval was between the highest dose where 4 mice survived and the lowest dose where 4 mice died within 5 minutes after an i.v. bolus dose.

# e) Effect on heart rate in conscious rat

The animal is slightly anaestetized by ether and an infusion cannula is inserted into a tail vein. While still asleep the rat is placed in a simple device, made of a coarse, somewhat elastic net fixing the rat in a constant position. Electrodes are attached to the extremities and connected to an ECG-pulse pre-amplifier and a Grass polygraph. By recording the ECG, the heart rate can then be determined.

Before any substance is given the animal has regained consciousness and the heart rate has been constant for at least 15 minutes.

The substance is injected, i.v. in the infusion cannula and flushed with physiological saline.

ECG is recorded 0.25, 0.5, 1, 2, 3 and 5 minutes after completed injection and then every 5 minutes until the original heart rate is obtained.

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Substance	Antichol. effect IC <sub>50</sub> (M)	Anti-N.A. effect IC <sub>50</sub> (M)	Anti-Ca effect IC <sub>50</sub> (M)	Anti-HI effect IC <sub>50</sub> (M)	Acute toxicity i.v. mg/kg	Lethal dose mg/kg	Effect on heart rate threshold dose mg/kg
CH <sub>3</sub> H CH-CH <sub>2</sub> -CH-N C(CH <sub>3</sub> ) <sub>3</sub> Terodiline (prior art)	5.2×10 <sup>-7</sup>	2.4×10 <sup>-6</sup>	10-5	4×10 <sup>-6</sup>	15-20	20	1-3
CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH <sub>3</sub> CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH <sub>3</sub> GB-A-1.169.944 (antidepressant)	1.2×10 <sup>-6</sup>	4.4×10 <sup>-6</sup>	2.1x10 <sup>-5</sup>	3.7×10 <sup>-7</sup>	10-15	15	
CH(CH <sub>3</sub> ) <sub>2</sub> CH-CH <sub>2</sub> -CH <sub>2</sub> -N  Racemate	1.8×10-8	01	1.5×10 <sup>-5</sup>	7×10 <sup>-6</sup>	10-20	20	[-3
la (+)-isomer of 1	1.8×10 <sup>-8</sup>						
1b (-)-isomer of 1	1.4×10 <sup>-8</sup>						
2 C(CH <sub>3</sub> ) <sub>3</sub> C(CH <sub>3</sub> ) <sub>3</sub>	1.5x10 <sup>-7</sup>	3.5×10 <sup>-6</sup>	9×10 <sup>-6</sup>		10-20	20	
3 CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH(CH <sub>3</sub> ) <sub>2</sub>	2.4×10 <sup>-7</sup>	3.6×10 <sup>-6</sup>	<sup>+-01</sup>		3-10	10	

Table I (cont.)

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Effect on heart rate threshold dose mg/kg				5-1		
Lethal dose mg/kg	0 †	20	20	45	>20	50
Acute toxicity i.v. mg/kg	30-40	10-20	10-20	30-45	> 20	30-50
Anti-III effect IC <sub>50</sub> (M)	10-5			10-5	1.3×10 <sup>-5</sup>	3×10 <sup>-6</sup>
Anti-Ca effect IC <sub>50</sub> (M)	6×10 <sup>-6</sup>	6.5x10 <sup>-6</sup>	9-01x9	3×10 <sup>-5</sup>	6.5×10 <sup>-5</sup>	6.5×10 <sup>-5</sup>
Anti-N.A. effect IC <sub>50</sub> (M)	5.5×10 <sup>-6</sup>		- 1	3.8×10 <sup>-5</sup>	3×10 <sup>-5</sup>	5×10 <sup>-5</sup>
Antichol. effect IC <sub>50</sub> (M)	1.5×10 <sup>-8</sup>	1.3x10 <sup>-8</sup>	1.3×10 <sup>-6</sup>	4.9×10 <sup>-9</sup>	2.0×10 <sup>-7</sup>	1.9×10 <sup>-8</sup>
Substance	4 CH(CH <sub>3</sub> ) <sub>2</sub> H <sub>3</sub> C CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH(CH <sub>3</sub> ) <sub>2</sub>	4a. (+)-isomer of 4 tartrate	4b. (-)-isomer of 4 tartrate	C(CH <sub>3</sub> ) <sub>3</sub> CH <sub>2</sub> -CH <sub>2</sub> -CH <sub>2</sub> -N CH <sub>3</sub> OH	HOCHON 6 CH-CH2-CH2-N CH3	7 ( ) CH(CH <sub>3</sub> ) <sub>2</sub> HO ( ) CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH(CH <sub>3</sub> ) <sub>2</sub>

Table I (cont.)

Substance	Antichol. effect IC <sub>50</sub> (M)	Anti-N.A. effect IC <sub>50</sub> (M)	Anti-Ca effect IC <sub>50</sub> (M)	Anti-HI effect IC <sub>50</sub> (M)	Acute toxicity i.v. mg/kg	Lethal dose mg/kg	Effect on heart rate threshold dose mg/kg
8 Cl OH CCH <sub>2</sub> -N CCCH <sub>3</sub> )3	3.1×10 <sup>-8</sup>	5×10 <sup>-5</sup>	>5x10 <sup>-5</sup>	7×10 <sup>-6</sup>	9 <	9 ^	
9 CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH <sub>3</sub>	1.6×10 <sup>-8</sup>	5×10 <sup>-5</sup>	2.5×10 <sup>-5</sup>	1.2×10 <sup>-6</sup>		20	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	6.2×10 <sup>-8</sup>	4×10-6	-7×10-6	2.5x10 <sup>-6</sup>		ن	
H <sub>3</sub> C. OH CH <sub>2</sub> -CH <sub>2</sub> -N CH(CH <sub>3</sub> ) <sub>2</sub>	1.0x10 <sup>-8</sup>	5.5x10 <sup>-6</sup>	10-5	2.5×10 <sup>-6</sup>	10-20	20	
12 HO CH-CH <sub>2</sub> -CH <sub>2</sub> -N CH <sub>3</sub>	4.7×10 <sup>-7</sup>		2.3×10 <sup>-5</sup>	8.0×10 <sup>-6</sup>	15-30	30	
13 (OH CH2-CH2-N CH(CH3)2 OH CH3)2	6-01×0-6	3×10 <sup>-5</sup>	1.5x10 <sup>-5</sup>	2×10 <sup>-5</sup>	5-10	. 10	·

### Example A

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### Preparation of tablets

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70		Ingredients	mg/tablet
	· 1.	Compound 1 in Table 1	2.0
15	2.	Cellulose, microcrystal- line	57.0
20	3.	Calcium hydrogen phosphate	15.0
	4.	Sodium starch glycolate	5.0
	5.	Silicon dioxide, colloidal	0.25
25	6.	Magnesium stearate	0.75
			80.0 mg

The compound 1 according to the invention is mixed with ingredients 2, 3, 4 and 5 for about 10 minutes. The magnesium stearate is then added, the resultant mixture being mixed for about 5 minutes and then compressed into tablet form with or without filmcoating.

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### Example B

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#### Preparation of capsules

		Ingredients	mg/capsule
45	1.	Compound 1 in Table 1	2
	2.	Lactose	186
	3.	Corn starch	20
	4.	Talc	15
50	5.	Magnesium	2
		stearate	
			225 mg

55 The compound 1 according to the invention is mixed with ingredients 2 and 3 and then milled. The resulting mixture is then mixed with ingredients 4 and 5 and then filled into capsules of appropriate size.

#### Claims

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1. 3,3-Diphenylpropylamines of formula I

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wherein R<sup>1</sup> signifies hydrogen or methyl, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> independently signify hydrogen, methyl, methoxy, hydroxy, carbamoyl, sulphanoyl or halogen, and X represents a tertiary amino group of formula II

wherein R<sup>5</sup> and R<sup>6</sup> signify non-aromatic hydrocarbyl groups, which may be the same or different and which together contain at least three carbon atoms, and wherein R<sup>5</sup> and R<sup>6</sup> may form a ring together with the amine nitrogen,

their salts with physiologically acceptable acids and, when the compounds can be in the form of optical isomers, the racemic mixture and the individual enantiomers.

2. 3,3-Diphenylpropylamines according to claim 1, wherein each of  $R^5$  and  $R^6$  independently signifies a saturated hydrocarbyl group, especially saturated alifatic hydrocarbyl groups such as  $C_{1-8}$ -alkyl, especially  $C_{1-6}$ -alkyl, or adamantyl,  $R^5$  and  $R^6$  together comprising at least three, preferably at least four carbon atoms.

3. 3,3-Diphenylpropylamines according to claim 1 or 2, wherein R<sup>5</sup> and R<sup>6</sup> taken together form a ring with the amine nitrogen.

4. 3,3-Diphenylpropylamines according to claim 1, 2 or 3, wherein R<sup>5</sup> and/or R<sup>6</sup> carries at least one hydroxy substitutent:

5. 3,3-Diphenylpropylamines according to any one of the preceeding claims, wherein at least one of R<sup>5</sup> and R<sup>6</sup> comprises a branched carbon chain.

6. 3,3-Diphenylpropylamines according to any one of claims 1-5, wherein X signifies any of the following groups a) - f), each of which may carry at least one hydroxy substituent:

a) 
$$-N \stackrel{\text{CH(CH}_3)_2}{\text{CH(CH}_3)_2}$$
, b)  $-N \stackrel{\text{CH}_3}{\text{C(CH}_3)_3}$ , c)  $-N \stackrel{\text{CH}_3}{\text{C(CH}_3)_2} \text{CH}_2 \text{CH}_3$ ,

d) 
$$\stackrel{CH_3}{\overset{CH_3}{\overset{CH_3}{\overset{CH_2}{\overset{CH_2}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}{\overset{CH_2}{\overset{CH_2}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}{\overset{CH_3}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{CH_3}}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{CH_3}}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{CH_3}}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{CH_3}}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}{\overset{CH_3}}}{\overset{C}}{\overset{CH_3}}{\overset{C}}{\overset{C}}}{\overset{C}}}{\overset{C}}{\overset{C}}{\overset{C}}{\overset{C}}}{\overset{C}}}{\overset{C}}{\overset{C}}$$

7. 3,3-Diphenylpropylamines according to claim 1, selected from the group consisting of the following compounds, their salts with physiologically acceptable acids and, where possible, their racemates and individual enantiomers:

N,N-diisopropyl-3-(2-hydroxy-5-methylphenyl)-3-phenylpropylamine,

N-methyl-N-tert.butyl-3-(2-hydroxyphenyl)-3-phenylpropylamine, N-methyl-N-tert.butyl-3-(2,4-dihydroxyphenyl)-3-phenylpropylamine, N-methyl-N-tert.butyl-3,3-bis-(2-hydroxyphenyl)propylamine, N,N-diisopropyl-3,3-bis-(2-hydroxyphenyl)propylamine, N,N-diisopropyl-3-(2,5-dihydroxyphenyl)-3-phenylpropylamine, N-methyl-N-tert.butyl-3-(2,5-dihydroxyphenyl)-3-phenylpropylamine, N,N-diisopropyl-3-(2-methoxyphenyl)-3-phenylpropylamine, N-[3-(2-methoxyphenyl)-3-phenylpropyl]-2,2,6,6-tetramethylpiperidine,

(+)-N,N-diisopropyl-3-(2-hydroxy-5-methylphenyl)-3-phenylpropylamine.
8. 3,3-Diphenylpropylamines according to any one of claims 1-7 for use as pharmaceutically active

substances, especially as anticholinergic agents.

9. A pharmaceutical composition comprising a 3,3-diphenylpropylamine according to any one of claims 1-7 and a compatible pharmaceutical carrier.

10. Use of a 3,3-diphenylpropylamine according to any one of claims 1-7 for preparing an anticholinergic

11. A method for preparing 3,3-diphenylpropylamines according to any one of claims 1-7, comprising:

a) reacting a reactively esterified 3,3-diphenylpropanol of formula III

$$\begin{array}{c}
R^{2} \\
\hline
O-OR^{1} \\
CH-CH_{2}-CH_{2}-Y
\end{array}$$
III

wherein  $R^1$ - $R^4$  are as defined above, any hydroxy groups may be protected and Y is a leaving group, with an amine of formula IV

H-X IV

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wherein X is as defined above, or

b) reducing a 3,3-diphenylpropionamide of formula V

wherein R1-R4 and X are as defined above and any hydroxy groups may be protected, or c) N-methylating a secondary 3,3-diphenylpropylamine VI

wherein R1-R4 are as defined above and any hydroxy groups may be protected, and wherein Z has the same meaning as R5 and R6 with the exception of methyl, or d) reducing a 3,3-diphenylpropylamine of formula VIIa or VIIb

VIIa

wherein R1-R4 and X are as defined above and any hydroxy groups may be protected, and W signifies

a hydroxy group or a halogen atom, and i) when necessary splitting off hydroxy protecting groups in the compounds obtained, if desired after

mono or di-halogenation of one or both of the phenyl rings, and/or ii) if desired converting obtained bases of formula I into salts thereof with physiologically acceptable acids, or vice versa, and/or

iii) if desired separating an obtained mixture of optical isomers into the individual enantiomers, and/or

iv) if desired methylating an ortho-hydroxy group in an obtained compound of formula I, wherein  $R^1$  is hydrogen and/or  $R^4$  is hydroxy.



PARTIAL EUROPEAN SEARCH REPORT which under Rule 45 of the European Patent Convention shall be considered, for the purposes of subsequent proceedings, as the European search report

Application number EP 89850017.8

	DOCUMENTS CON			
Category	Citation of document of re	with indication, where appropriate, levant passages	Relevant to claim	CLASSIFICATION OF TH APPLICATION (Int. CI.4
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EP 89850017.8

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